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(\$4) Title: TEMPLATED MOLECULES AND MITHODS FOR USING SUCH MOLECULES

Chemical Display - Principle.

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molecules. In one aspect of the invention, the templated molecules are linked to the template which templated the synthesis thereof. The intion allows the generation of (57) Abstract: The present invention relates to a method for synthesising templated libraries which can be screened for e.g. therapeutic activity.

WO 02/103008 A2 3 DK-3460 Birkerød (DK). ANDERS GODSKESEN, Michael [DK/DK]; Plantagekrogen 8, DK-2950 Vedbæk (DK). SCHRØDER GLAD, Sanne

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without international search report and to be republished upon receipt of that report For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

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A method for synthesising a templated molecule

Technical Field of the Invention

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Biological systems allow template-directed synthesis of α-peptides. The present invention enables a system that allows template-directed synthesis of other types of polymers as well as α-peptides. The present invention relates to templated molecules and templated molecules linked to a predetermined template. The templated molecules comprise a sequence of functional groups that are linked together. Each functional group is initially linked to an element capable of complementing a predetermined coding element of the template. Following complementation of a coding element, or complementation of a plurality of coding element, the appended functional groups are linked and the templated moleculed is formed.

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15 Background

The central dogma in biology describes the flow of information as a one-way process from DNA to RNA to polypeptide. Accordingly, DNA is transcribed by a RNA polymerase into mRNA; and the mRNA is subsequently then translated into protein by the ribosomes and fRNAs.

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The direct relation between the DNA and the protein, I.e., the fact that the sequence of triplet codons defines the sequence of α -amino acid residues in a polypeptide, has allowed the development of numerous molecular biological methods, in which the experimenter manipulates the DNA (mutagenizes, recombines, deletes, inserts, etc), and then uses in vivo systems (e.g., microbes) or in vitro systems (e.g., Zubay in vitro expression systems) to transfer the resulting changes from the DNA level to the level of the templated polypeptide, i.e., to mutate, recombine, delete, insert, etc. the polypeptide.

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Several systems have been invented that allows a flow of information from polypeptide to DNA. These systems are phage display, ribosome/polysome display, peptides-on-plasmid display, and other systems. These systems introduce a physical link between the template (e.g., DNA) and the templated molecule (polypeptide). As a result, it is possible, from a population of templated molecules linked to the tem-

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plate that templated the synthesis of the molecule, to first enrich for a desired characteristic of the templated molecule (e.g., binding of the templated molecule to an affinity column), and then amplify the enriched population of templated molecules through amplification of its template (DNA or RNA), followed by translation of the amplified templates. These methods have been used to identify polypeptides with novel and/or improved features from libraries consisting of from a million to about 10¹⁵ polypeptides.

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The critical feature of the prior art systems is the amplifiability of the templated molecule, through amplification of its template. Thus, after the selection step in which molecules with the desired property are enriched, the enriched population may be amplified and then taken through yet a selection step, etc. - the process of selection-and-amplification may be repeated many times. In this way the "noise" of the selection assay is averaged out over several selection-and-amplification rounds, and even if the individual selection step only enriches e.g. 10-fold, a theoretical enrichment of 10¹² can be reached after 12 selection-and-amplification rounds. Had the molecules <u>not</u> been amplifiable, the same enrichment would have had to be achieved in a single screening step, which means that the enrichment in this one step would have had to be 10¹², and the assay should still have the same overall stringency (accuracy). This is practically impossible with current technologies.

In the field of chemistry, a different combinatorial approach has been developed.

This approach involved the parallel synthesis of millions of related compounds, in an array (where each position defined a specific compound), or on beads (where one bead carried many copies of the same compound). The population of compounds were then screened for desired characteristics. Importantly, this type of combinatorial library has no means for amplification, and therefore requires the use of very stringent screening methods, as explained above. Recently, the trend in for example medicinal chemistry has therefore been to use less diverse, but better character-

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Principles for tagging chemical libraries have also been developed. For example, systems that employed DNA oligos to tag molecule libraries have been developed as exemplified herein below. The tag is used as a means of identification, but can-

zed libraries.

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not be used to template the synthesis of the tagged molecule. Therefore, despite the tag, these systems still require a very efficient screening method.

The below listed references illustrate some of the above-mentioned short-comings of the prior art methods in the field of the invention.

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EP 0 604 552 B1 relates to a method for synthesizing diverse collections of oligomers. The invention involves the use of an identifier tag to identify the sequence of monomers in an oligomer. The identifier tags facilitate subsequent identification of reactions through which members of a library of different synthetic compounds have been synthesised in a component by component fashion.

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EP 0 643 778 B1 relates to encoded combinatorial chemical libraries. Each of a collection of polypeptides is labelled by an appended "genetic" tag, itself constructed by chemical synthesis, to provide a "retro-genetic" way of specifying each polypeptide.

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EP 0 773 227 At relates to a method for preparing a new pharmaceutical drug or diagnostic reagent, which includes the step of screening, against a ligand or receptor, a library of different synthetic compounds obtainable by synthesis in a component by component fashlon.

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US 4,863,857 relates to a method for determining the amino acid sequence of a polypeptide complementary to at least a portion of an original peptide or protein. In one aspect the method involves: (a) determining a first nucleotide sequence of a first nucleic acid coding for the blosynthesis of at least a portion of the original peptide or protein; (b) ascertaining a second nucleotide sequence of a second nucleic acid, which base-pairs with the first nucleotide sequence of the first nucleic acid, the first and second nucleic acids pairing in antiparallel directions; and (c) determining the amino acid sequence of the complementary polypeptide by the second nucleotide sequence when read in the same reading frame as the first nucleotide sequence.

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US 5,162,218 relates to polypeptide compositions having a binding site specific for a particular target ligand and further having an active functionality proximate the binding site. The active functionality may be a reporter molecule, in which case the

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polypeptide compositions are useful in performing assays for the target ligand. Also disclosed are methods for preparing polypeptides having active functionalities proximate their binding site, said method comprising the step of combining the polypeptide specific for the target ligand with an affinity label having a reactive group attached thereto. The reactive group is then covalently attached to an amino acid side chaln proximate the binding site and cleaved from the substrate. The substrate

is subsequently eluted, leaving a moiety of the reactive group covalently attached to

the polypeptide. The active funtionality may then be attached to the molety.

US 5,270,170 relates to a random peptide library constructed by transforming host cells with a collection of recombinant vectors that encode a fusion protein comprised of a DNA binding protein and a random peptide and also encode a binding site for the DNA binding protein. The fusion protein can be used for screening ligands. The screening method results in the formation of a complex comprising the fusion protein bound to a receptor through the random peptide ligand and to the recombinant DNA vector through the DNA binding protein.

US 5,539,082 relates to a novel class of compounds, known as peptide nucleic acids capable of binding complementary ssDNA and RNA strands more strongly than a corresponding DNA. The peptide nucleic acids generally comprise ligands such as naturally occurring DNA bases attached to a peptide backbone through a suitable linker.

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US 5,574,141 relates to functionalized carrier materials for the simultaneous synthesis and direct labeling of oligonucleotides as primers for template-dependent enzymatic nucleic acid syntheses. The polymeric carriers are loaded with nucleic acid building blocks which In turn contain labelling groups or precursors thereof. The polymeric carrier loaded in this way serves as a solid or liquid phase for the assembly of oligonucleotides which can be used as primers for a template-dependent enzymatic nucleic acid synthesis such as in sequencing analysis or in the polymerase chain reaction (PCR).

US 5,573,905 relates to an encoded combinatorial chemical library comprised of a plurality of bifunctional molecules having both a chemical polymer and an identifier oligonucleotide sequence that defines the structure of the chemical polymer. Also

library to identify chemical structures within the library that bind to biologically active described are the bifunctional molecules of the library, and methods of using the molecules in preselected binding interactions.

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identification and discovery of agents which are inhibitors and activators of RNA and able primer motecutes, and any necessary accessory motecutes, catalytic extension the incorporation of a functional polymerase binding site sequence (PBS) into a nucleic acid molecule which is chosen for its ability to confer a discernible characteris-US 5,597,697 relates to a screening assay for inhibitors and activators of RNA and DNA-dependent nucleic acid polymerases. The essential feature of the invention is template directed nucleic acid polymerase. In the presence of the polymerase, suit-DNA-dependent nucleic acid polymerases. The invention provides methods for the partial or total elimination of (or increase in) the characteristic conferring activity of tic via its sequence specific activity such that the incorporation of the PBS renders of the strand of nucleic acids complementary to the template occurs, resulting in a the reporter-template molecule described due to the antisense effects of the comthe nucleic acid molecule a functional template for a predetermined RNA or DNAplementary strand or other polymerase-mediated effects.

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US 5,639,603 relates to a method for synthesizing and screening molecular diversity by means of a general stochastic method for synthesizing compounds. The method can be used to generate large collections of tagged compounds that can be screened to identify and isolate compounds with useful properties.

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tor are isolated and their sequence of base moieties is determined. Also disclosed is US 5,698,685 relates to a morpholino-subunit combinatorial library and a method for generating a compound capable of interacting specifically with a selected macromolecular ligand. The method involves contacting the ligand with a combinatorial library non-nucleobase side chains. Oligomer molecules that bind specifically to the recep of oligomers composed of morpholino subunits with a variety of nucleobase and a combinatorial library of oligomers useful in the method and novel morpholinosubunit polymer compositions.

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compounds by means of a general stochastic method for synthesizing random oli-US 5,708,153 relates to a method for synthesizing diverse collections of tagged 33

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tion tags on the particles to facilitate identification of the sequence of the monomers gomers on particles. A further aspect of the invention relates to the use of identificein the oligomer.

- solubility. The peptide nucleic aclds comprise ligands selected from a group consistids, which bind complementary DNA and RNA strands more strongly than the corre-US 5,719,262 relates to a novel class of compounds, known as peptide nucleic acng of naturally-occurring nucleobases and non-naturally-occurring nucleobases sponding DNA or RNA strands, and exhibit increased sequence specificity and attached to a polyamide backbone, and contain alkylamine side chains. ß 2
- binding affinity, where the products may be detached from the particle or retained on he particle. The reaction history of the particles which are positive for the characteristic can be determined by the release of the tags and analysis to define the reaction able tags. The particles may be screened for a characteristic of interest, particularly and stage, as the same or different bit of information. Various products can be probinary or higher code, so as to define a plurality of choices with only a few detachduced in the multi-stage synthesis, such as oligomers and synthetic non-repetitive organic molecules. Particularly, pluralities of identifiers may be used to provide a ags. Encoded combinatorial chemistry is provided, whereby sequential synthetic schemes are recorded using organic molecules, which define choice of reactant, JS 5,721,099 relates to encoded combinatorial chemical libraries encoded with listory of the particle. 5 ೪
- ibrary to Identify chemical structures within the library that bind to biologically active US 5,723,598 relates to an encoded combinatorial chemical library comprised of a plurality of bifunctional molecules having both a chemical polymer and an identifier oligonucleotide sequence that defines the structure of the chemical polymer. Also described are the bifunctional molecules of the library, and methods of using the molecules in preselected binding interactions.

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JS 5,770,358 relates to tagged synthetic oligomer libraries and a general stochastic method for synthesizing random oligomers. The method can be used to synthesize compounds to screen for desired properties. The use of identification tags on the oligomers facilitates Identification of oligomers with desired properties

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US 5,786,461 relates to peptide nucleic acids having amlno acid side chains. A novel class of compounds, known as peptide nucleic acids, bind complementary DNA and RNA strands more strongly than the corresponding DNA or RNA strands, and exhibit increased sequence specificity and solubility. The peptide nucleic acids comprise ligands selected from a group consisting of naturally-occurring nucleobases and non-naturally-occurring nucleobases and contain alkylamine side chains.

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US 5,789,162 relates to a method for synthesizing diverse collections of oligomers. A general stochastic method for synthesizing random oligomers on particles is disclosed. A further aspect of the invention relates to the use of identification tags on the particles to facilitate identification of the sequence of the monomers in the oll-gomer.

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US 5,840,485 relates to topologically segregated, encoded solid phase libraries. Libraries of synthetic test compounds are attached to separate phase synthesis supports that also contain coding molecules that encode the structure of the synthetic test compound. The molecules may be polymers or multiple nonpolymeric molecules. The synthetic test compound can have backbone structures with linkages such as amide, urea, carbamate (i.e., urethane), ester, amino, sulfide, disulfide, or carbon-carbon, such as alkane and alkene, or any combination thereof. The synthetic test compound can also be molecular scaffolds, or other structures capable of acting as a scaffolding. The invention also relates to methods of synthesizing such libraries and the use of such libraries to identify and characterize molecules of interest from among the library of synthetic test compounds.

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US 5,843,701 relates to systematic polypeptide evolution by reverse translation and a method for preparing polypeptide ligands of target molecules wherein candidate mixtures comprised of ribosome complexes or mRNA:polypeptide copolymers are partitioned relative to their affinity to the target and amplified to create a new candidate mixture enriched in ribosome complexes or mRNA:polypeptide copolymers with an affinity to the target.

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US 5,846,839 relates to a method for hard-tagging an encoded synthetic library. Disclosed are chemical encryption methods for determining the structure of compounds formed in situ on solid supports by the use of specific amine tags which, after compound synthesis, can be deencrypted to provide the structure of the compound found on the support.

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US 5,922,545 relates to methods and compositions for identifying peptides and single-chain antibodies that bind to predetermined receptors or epitopes. Such peptides and antibodies are identified by methods for affinity screening of polysomes displaying nascent peptides.

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US 5,958,703 relates to methods for screening libraries of complexes for compounds having a desired property such as the capacity to bind to a cellular receptor. The complexes in such libraries comprise a compound under test, a tag recording at least one step in synthesis of the compound, and a tether susceptible to modification by a reporter molecule. Modification of the tether is used to signify that a complex contains a compound having a desired property. The tag can be decoded to reveal at least one step in the synthesis of such a compound

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20 US 5,986,053 relates peptide nucleic acid complexes of two peptide nucleic acid strands and one nucleic acid strand. Peptide nucleic acids and analogues of peptide nucleic acids are used to form duplex, triplex, and other structures with nucleic acids and to modify nucleic acids. The peptide nucleic acids and analogues thereof also are used to modulate protein activity through, for example, transcription arrest, tran-

25 scription initiation, and site specific cleavage of nucleic acids.

US 5,998,140 relates to methods and compositions for forming complexes intraceltularly between dsDNA and oligomers of heterocycles, aliphatic amino acids, particularly omega-amino acids, and a polar end group. By appropriate choice of target
sequences and composition of the oligomers, complexes are obtained with low dissociation constants.

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US 6,060,596 relates to an an encoded combinatorial chemical library comprised of a plurality of bifunctional molecules having both a chemical polymer and an Identifier oligonucleotide sequence that defines the structure of the chemical polymer. Also

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described are the bifunctional molecules of the library, and methods of using the library to identify chemical structures within the library that bind to biologically active molecules in preselected binding interactions.

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US 6,080,826 relates to Template-directed ring-closing metathesis and ring-opening metathesis polymerization of functionalized dienes. Functionalized cyclic olefins and methods for making the same are disclosed. Methods include template-directed ring-closing metathesis ("RCM") of functionalized acyclic dienes and template-directed depolymerization of functionalized polymers possessing regularly spaced sites of unsaturation. Although the template species may be any anion, cation, or dipolar compound, cationic species, especially alkall metals, are preferred. Functionalized polymers with regularly spaced sites of unsaturation and methods for making the same are also disclosed. One method for synthesizing these polymers is by ring-opening metathesis polymerization ("ROMP") of functionalized cyclic olefins.

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US 6,127,154 relates to compounds which possess a complementary structure to a desired molecule, such as a biomolecule, in particular polymeric or oligomeric compounds, which are useful as in vivo or in vitro diagnostic and therapeutic agents are provided. Also, various methods for producing such compounds are provided.

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US 6,140,493 relates to a method for synthesizing diverse collections of oligomers. A general stochastic method for synthesizing random oligomers is disclosed and can be used to synthesize compounds to screen for desired properties, Identification tags on the oligomers facilitates Identification of oligomers with desired properties.

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US 6,140,496 relates to building blocks for preparing oligonucleotides carrying nonstandard nucleobases that can pair with complementary non-standard nucleobases so as to fit the Watson-Crick geometry. The resulting base pair joins a monocyclic six membered ring paining with a fused bicyclic heterocyclic ring system composed of a five member ring fused with a six member ring, with the orientation of the heterocycles with respect to each other and with respect to the backbone chain analogous to that found in DNA and RNA, but with a pattern of hydrogen bonds holding the base pair together different from that found in the AT and GC base pairs (a "nonstandard base pair").

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US 6,143,497 relates to a method for synthesizing diverse collections of random oligomers on particles by means of a general stochastic method. Also disclosed are identification tags located on the particles and used to facilitate identification of the sequence of the monomers in the oligomer.

US 6,165,717 relates to a general stochastic method for synthesizing random oilgomers on particles. Also disclosed are Identification tags located on the particles to facilitate identification of the sequence of the monomers in the oligomer.

10 US 6,175,001 relates to functionalized pyrimidine nucleosides and nucleotides and DNA's incorporating same. The modified pyrimidine nucleotides are derivatized at C5 to contain a functional group that mimics the property of a naturally occurring amino acid residues. DNA molecules containing the modified nucleotides are also provided.

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US 6,194,550 B1 relates to systematic polypeptide evolution by reverse translation, in particular a method for preparing polypeptide ligands of target molecules wherein candidate mixtures comprised of ribosome complexes or mRNA:polypeptide copolymers are partitioned relative to their affinity to the target and amplified to create a new candidate mixture enriched in ribosome complexes or mRNA;polypeptide copolymers with an affinity to the target.

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US 6,207,446 B1 relates to methods and reagents for the selection of protein molecules that make use of RNA-protein fusions.

US 6,214,553 B1 relates to methods and reagents for the selection of protein molecules that make use of RNA-protein fusions.

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WO 91/05058 relates to a method for the cell-free synthesis and isolation of novel genes and polypeptides. An expression unit is constructed onto which semi-random nucleotide sequences are attached. The semi-random nucleotide sequences are first transcribed to produce RNA, and then translated under conditions such that polysomes are produced. Polysomes which bind to a substance of interest are then isolated and disrupted; and the released mRNA is recovered. The mRNA is used to construct cDNA which is expressed to produce novel polypeptides.

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cules wherein candidate mixtures comprised of ribosome complexes or

WO 92/02536 relates to a method for preparing polypeptide ligands of target mole-

mRNA;polypeptIde copolymers are partitioned relative to their affinity to the target

and emplified to create a new candidate mixture enriched in ribosome complexes or

mRNA:polypeptide copolymers with an affinity to the target.

WO 93/03172 relates to a method for preparing polypeptide ligands of target molecules wherein candidate mixtures comprised of ribosome complexes or

10 mRNA:polypeptide copolymers are partitioned relative to their affinity to the target

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and amplified to create a new candidate mixture enriched in ribosome complexes or

mRNA:polypeptide copolymers with an affinity to the target.

WO 93/06121 relates to a general stochastic method for synthesizing random oil15 gomers on particles. Also disclosed are identification tags located on the particles to
facilitate identification of the sequence of the monomers in the oligomer.

WO 00/47775 relates to a method for generating RNA-protein fusions involving a

high-salt post-translational step.

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Additional references of relevance for present invention includes Bain et al. Nature, vol. 356, 1992, 537-539; Barbas et al. Chem. Int. Ed. vol. 37, 1998. 2872-2875;

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Summary of the Invention

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The present invention solves in a general way the above-mentioned problems and short-comings of the prior art. The invention relates to a system for templating molecules in general, such as polymers, and the template enables templated synthesis of the polymers, allowing in preferred embodiments amplification of the polymer. The

system therefore has the same overall characteristics as the natural system (information flow from template to templated molecule), as well as the characteristics of the recently invented ribosome-mediated systems (e.g., phage display), namely the physical link between template and templated molecule. However, the present invention does not involve ribosomes or tRNAs, and therefore allows templating of a

30 wide array of different polymers, including polymers that cannot be synthesised in a natural system based on ribosome-mediated translation of nucleic acids.

The templating process of the invention has significant advantages over the prior art. As the amplification of the recovered molecules (i.e., their templates) can be done by a parallel process in which all the recovered templates are present in the

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sams compartment (e.g., reagent tube or microtiter-plats well), and where the molecopying the molecules one-at-a-time, i.e., to "amplify" the molecules in a serial proccovery after a first selection round Involves e.g. 1010 different molecules, when the the Individual molecules is necessary. This is a huge advantage since a typical recules are proportionately amplified, no human intervention such as sequencing of starting material is a library of e.g. 1015 molecules. When working with such high numbers of molecules, it is practically impossible to "amplify" 10^{10} molecules by ess.

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The present invention generally relates to templated molecules and complexes comprising such molecules linked to a template that has directed the templatemolecules and the complexes are obtainable according to the methods of the directed synthesis of the templated molecule. In one aspect, the templated present invention.

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from building blocks comprising a functional entity comprising a functional group and complexes to a target species. The templated molecules are preferably synthesised complementing element by a cleavable linker, or a selectively cleavable linker. The element of the template, thus ensuring a one-to-one relationship between a coding emplated molecule. The functional entity of a building block is separated from a complementing element is capable of complementing a predetermined coding element - or a complementing element - and a functional entity, or a functional The present invention also discloses methods for synthesizing such templated reactive group capable of covalently linking functional groups and forming a molecules and/or complexes, methods for targeting such molecules and/or

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Also disclosed are methods for identifying the sequence of functional groups of a templated molecule, as well as methods for therapy and diagnostic methods

exploiting the templated molecules according to the invention. 8

either I) exclusively a-emino acids, or II) substantially exclusively naturally occurring ribonucleic acids. Also, when the templated molecules are peptides comprising The methods of the invention do not involve ribosome mediated translation of amino acids, such as at least 80 percent, for example 90 percent, such as 95

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percent, naturally occurring amino acids, the template does not comprise or essentially consist of a ribonucleic acid. A template denotes a sequence of coding elements, wherein each coding element is sequence of complementing elements, wherein each complementing element is linked to a neighbouring coding element. A complementing template denotes a inked to neighbouring complementing element. ß

capable of being linked - without forming part of the complementing template itself complementation of a plurality of coding elements by a plurality of complementing element. Accordingly, in one preferred embodiment, the functional group does not slements, each complementing element will define an appended functional group Following complementation of a coding element by a complementing element, or to a neighbouring functional group defined by a neighbouring complementing 9

participate in the complementation of a coding element in so far as no direct reaction Interaction - covalent or non-covalent - between the functional group and the coding or hybridization takes place between the coding element and the functional group. The term "reaction" means any reactive contact that results in the formation of an element. In another embodiment, the functional group of a templated molecule 5

forms part of the complementing template. ຂ

element of a template, and as each coding element in turn defines a predetermined functional group, the sequence of coding elements of the template will template the As each complementing element is capable of recognising a predetermined coding

synthesis of the templated molecule comprising a predetermined sequence of covalently linked functional groups. 22

According to preferred embodiments of the present invention, it is possible

i) to link a templated molecule comprising a plurality of functional groups to the emplate that templated the synthesis of the templated molecule. ജ

ii) to link neighbouring functional groups simultaneously with the complementation of nelghbouring coding elements by complementing elements defining said functional

groups,

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iii) to link neighbouring functional groups after the complementation of neighbouring coding elements by complementing elements defining said functional groups,

- iv) to link neighbouring furictional groups simultaneously with the formation of a complementing template, 2
- v) to link neighbouring functional groups after the formation of a complementing template,

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- complementing template without cleaving links between functional groups of a vi) to cleave one or more links between complementing elements of a templated motecule, and vice versa, and
- vii) to cleave the at least one linker separating the at least one functional entity from the at least one complementing element of a building block without cleaving the complementing template, र्घ
- viii) to cleave the at least one linker separating the at least one functional entity from the at least one complementing element of a building block without cleaving the link between the functional groups of the templated molecule, and 8
- complementing template and without cleaving the link between the functional groups ix) to cleave the at least one linker separating the at least one functional entity from the at least one complementing element of a building block without cleaving the of the templated molecule. 23

conditions wherein a selectively cleavable linker is not cleavable. Accordingly, it is functional groups of a templated molecule. Cleavable linkers are cleavable under Provided that complementation of neighbouring coding elements is achieved, the cleavable linker separating the functional entity from the complementing element neighbouring, functional groups of the templated molecule are capable of being linked irrespective of whether a complementing template is formed. Also, it is defining said functional entity without cleaving the link between neighbouring possible to link neighbouring functional groups and subsequently cleave the

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selectively cleavable linkers linking - in the same templated motecule - a subset of template-mediated synthesis of the templated molecule, wherein the template and complex comprising a templated molecule and the template that has directed the complementing elements and functional groups. It is thus possible to obtain a possible to cleave the cleavable linkers linking complementing elements and functional groups in a templated molecule without at the same time cleaving

- the templated molecule are linked by one or more, preferably one, selectively cleavable linker(s).
- possible using prior art "tags" generated by step-by-step synthesis. Accordingly, the The generation of additional templated molecules can be directed by the template without any need for sequencing or any other form of characterisation. This is not complexes of the invention comprising a templated molecule linked to a template nakes it possible to rapidly select and ampiify desirable, templated molecules.

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in a first aspect, the present invention provides a method for synthesising a templated molecule comprising a plurality of functional groups, said method comprising the steps of

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providing at least one template comprising a sequence of n coding elements,

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- wherein each coding element comprises at least one recognition group capable of recognising a predetermined complementing element, and

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wherein n is an integer of more than 1,

- providing a plurality of building blocks, wherein each building block comprises ≘
- recognition group capable of recognising a predetermined coding a) at least one complementing element comprising at least one element,

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 b) at least one functional entity comprising at least one functional group and at least one reactive group, and

 at least one linker separating the at least one functional entity from the at least one complementing element,

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- iii) contacting each of said coding elements with a complementing element capable of recognising said coding element,
- iv) optionally, obtaining a complementing element, and

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 obtaining a templated molecule comprising covalently linked, functional groups by linking, by means of a reaction involving reactive groups, a functional group of at least one functional entity to a functional group of another, functional entity,

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wherein the templated molecule is capable of being linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, and

wherein the synthesis of the templated molecule does not involve ribosome mediated-translation of a nucleic acid.

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In another aspect, the present invention relates to a templated molecule, a plurality of the same or different templated molecules, wherein preferably each of the templated molecules are obtainable by a method for synthesizing templated molecules according to the present Invention.

As the templated molecule and the template are separate entities capable of being linked by a single linker, the invention also relates to complexes comprising a templated molecule liked to the template that templated the synthesis of the templated molecule. The template capable of templating the synthesis of the templated molecule comprises either a sequence of coding elements, or a sequence

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of complementing elements, in which case the template is a complementing template. Accordingly, it is possible to cleave links between functional groups of a templated molecule without cleaving a complementing template or template that templated the synthesis of the templated molecule.

In another aspect there is provided a method for synthesising a complex comprising a templated molecule linked to the template that templated the synthesis of the templated molecule, wherein the templated molecule and the complex comprising the templated molecule linked to the template that templated the synthesis of the tem-

In further aspects of the invention there is provided a composition comprising a plurality of templated molecules, wherein each or at least some of the templated molecules are linked to the template that templated the synthesis of the templated molecule, in which case there is provided a plurality of complexes each comprising a templated molecule linked to the template that templated the synthesis of the templated molecule. The compositions may also comprise a templated molecule and unlinked thereto - the template that templated the synthesis of the templated molecule molecule.

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The amplifiability of the templated molecules of a library provides a library with a unique feature. This unique feature involves e.g. that a huge number of templated molecules can be screened by taking the library through repetitive processes of selection-and-amplification, in a parallel process where the library of molecules is treated as a whole, and where it is not necessary to characterise Individual molecules (or even the population of molecules) between selection-and-amplification rounds.

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10 It is possible according to various preferred embodiments of the invention to screen e.g. more than or about 10³ different templated molecules, such as more than or about 10⁴ different templated molecules, for example more than or about 10⁵ different templated molecules, such as more than or about 10⁴ different templated molecules, such as cules, for example more than or about 10⁴ different templated molecules, such as

35 more than or about 10⁸ different templated molecules, for example more than or

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about 10° different templated molecules, such as more than or about 10° different templated molecules, for example more than or about 10° different templated molecules, for example cules, such as more than or about 10° different templated molecules, for example more than or about 10° different templated molecules, such as more than or about 10° different templated molecules, for example more than or about 10° different templated molecules, such as more than or about 10° different templated molecules, such as more than or about 10° different templated molecules, such as more than or about 10° different templated molecules, such as more than or about 10° different templated molecules.

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As one may perform many repetitive rounds of parallel selection and parallel amplification processes, it is possible to enrich only e.g. 100 fold in each round, and still get a very efficient enrichment, of e.g. 10¹⁴ fold over a number of selection-and-amplification rounds (theoretically a 10¹⁴ fold enrichment is obtained after seven rounds each enriching 100 fold). To obtain a similar enrichment of 10¹⁴ fold using a non-amplifiable library, would require screenling conditions allowing 10¹⁴ fold enrichment in one "round" - and this is not practically possible using state-of-the-art screenling technologies. The templated molecules and/or the templates can furthermore be bound to a solid or semi-soild support.

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20 In even further aspects the methods of the invention - individually or as a combination - relates to a method for screening a composition of complaxes or templated molecules potentially having a predetermined activity,

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a method for assaying the predetermined activity potentially associated with the complexes or the templated molecules,

 a method for selecting complexes or templated molecules having a predetermined 30 activity, a method for amplifying the template that templated the synthesis of the templated molecule having, or potentially having a predetermined activity, and

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a method for amplifying the template that templated the synthesis of the templated molecule having, or potentially having, a predetermined activity, said method comprising the further step of obtaining the templated molecule in an at least two-fold increased amount.

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In yet another aspect there is provided a method for altering the sequence of a templated molecule, including generating a templated molecule comprising a novel or altered sequence of functional groups, wherein the method comprises the step of mutating the template that templated the synthesis of the original templated mole-

- cule. The method preferably comprises the steps of
- providing a first template capable of templating the first templated molecule, or a plurality of such templates capable of templating a plurality of first templated molecules,

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modifying the sequence of the first template, or the plurality or first templates, and generating a second template, or a plurality of second templates, wherein said second template(s) is capable of templating the synthesis of a second templated molecule, or a plurality of second templated molecules,

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wherein said second templated molecule(s) comprises a sequence of covalently linked, functional groups that is not identical to the sequence of functional groups of the first templated molecule(s), and optionally

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 iii) templating by means of said second template(s) a second templated molecule, or a plurality of such second templated molecules.

The above-mentioned method exploits that a templated synthesis (Figure 1)

30 in one embodiment involves a single-stranded, modifiable intermediate in the form of a template. In the case where this template comprises a nucleotide strand comprising deoxyribonucleotides or ribonucleotides, most molecular biological methods can be applied to modify the template, and therefore to modify the templated polymer.

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The below-mentioned list of molecular biological methods that can be applied to the templated polymers of this invention is therefore far from comprehensive, but merely serves to illustrate that almost any relevant molecular biological method can be applied to the templated polymers as a result of the present invention.

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In cases where nucleotides with non-natural bases are part of the template, some of the molecular biology methodologies may not be applicable. This will primarily depend on the substrate specificty of the enzymes involved (e.g., the Taq DNA polymenase in a PCR reaction; restriction enzyme in USE protocol; etc). Also, methods that involve an in vivo step (e.g., transformation of E. coli for amplification of plasmid DNA) may only have a limited feasibility for those nucleotides. Several nucleotides with non-natural bases are, however, known to be incorporated into oligonucleotides by several wildtype and mutant polymerases, and therefore, the use of nucleotides with non-natural bases does not seriously limit the number of in vitro molecular biology methods that can be applied to templated molecules.

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Table 1. Molecular Biology applicable to the templated polymers of this Invention

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In vivo and in vitro amplification, recombination and mutagenesis

 Kunkel site-directed mutagenesis, using one or mutiple (e.g., 50) different mutagenic oligos at below-saturating concentrations, i.e., generating a combinatorial library

USE (Unique Site-directed Elimination), using one or multiple (e.g., 50 different mulagenic oligos) at below-saturating concentrations, i.e., generating a combinatorial library

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PCR (Polymerase Chain Reaction)

LCR (Ligase Chain Reaction)

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 PCR shuffiling, including family shuffling (shuffling sequences containing blocks with particular homology), and directed shuffling where oligos are spiked into the reaction to direct the shuffling process in a certain direcOther types of shuffling, e.g. homologous recombination in yeast; shuffling protocols as developed at the companies Phylos, Energy Biosystems, Diversa and by Frances Amold.

Cassette mutagenesis

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 Other polymerase- or PCR-based methods, e.g., overlap extension, gene synthesis, and error-prone PCR

Chemical or UV-induced mutagenesis

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Wildtype or variant template synthesis and translation into templated
 polymer (wildtype in this respect means the template sequence that will
template the synthesis of the known ("wildtype") polymer, variant in this
respect means a partly randomised or spiked template sequence that will
template the synthesis of a variant of the known polymer)

Specific cleavage by restriction enzymes

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Ligation by DNA or RNA ligases; "gene splicing"

Affinity selections (using the template-templated polymer complex)

Sequending

 Arraying the polymers on "DNA chips", by using the template as a tag that binds a DNA array

instead of isolating the (underivatized) template strand, it may be desirable to apply

the molecular biological methods to either the template-complementing template

double-hellx or to the derivatized complementing template. The derivatized

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identifying a pharmaceutical agent, or a diagnostic agent, in the form of candidate plurality of drug targets with at least one predetermined, templated molecule, and templated molecules capable of Interacting with said drug targets. In yet another aspect there is provided a method for identifying a target, including a ligands or receptor moletles with at least one predetermined, templated, molecule, and Identifying drug targets in the form of ligands or receptor moleties capable of drug target, wherein sald method comprises the step of screening a plurality of interacting with said templated molecules. S

connected the functional entity and the complementing element. Many polymenases

template may at this point contain unpolymerized functional entities; polymerized

functional entities; or a trace left behind from the cleaving of the linker that

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specificity of the enzymes involved whether it will be feasible to use the derivatized

question. The skilled person will be capable of evaluating the feasibility of various

practical approaches in this respect.

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complementing template as a starting point for the molecular biological method in

and other enzymes are likely to be feasible using the (derivatized) complementing

template as starting point. It will primarily depend on the substrate- or template

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degree of derivatization. Therefore, many in vitro methods involving polymerases

and other enzymes are known to accept DNA- or RNA-templates with a high

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cell surfaces, solld or semi-solid surfaces, as well as any other physical or molecular having an affinity for a predetermined target, including a drug target, as well as to targets, including drug targets, in the form of ligands, receptor moieties, enzymes, The present Invention also relates to any isolated or purified templated molecule entity or surface having an affinity for a predetermined templated molecule.

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The present invention also pertains to building blocks used for synthesising the tem-

plated molecule and to complexes comprising such building blocks. In another as-

templated molecule comprises or essentially consists of a molecular entity capable

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of binding to another molecular entity in the form of a target moleculer entity or a

binding partner

molecule according to the invention. In a preferred embodiment of this aspect, the

pect there is provided the use of a building block for the synthesis of a templated

The templated molecule is preferably a medicament capable of being administered

idual and treating a clinical condition in said individual in need of such treatment.

In a pharmaceutically effective amount in a pharmaceutical composition to an indi-

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an individual in need thereof, said method comprises the step of administering to the of the present invention and having an affinity for a predetermined target, including a In even further aspects of the invention there is provided a method for treatment of individual a pharmaceutically effective amount of a molecule identified by a method

drug target.

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pharmaceutically effective amount of an isolated or purified ligand or recaptor molety need thereof, said method comprises the step of administering to the individual a In a still further aspect there is provided a method for treatment of an Individual in having an affinity for a predetermined templated molecule according to the inven-

tion. The isolated or purified ligand or receptor molety is preferably Identified by the above-mentioned method of identification of the invention. 22

coding element involves one or more polymerases or transcriptases. Thus, in accorthe building blocks may be a di- or oligonucleotides. While mononucleotides are the The present invention may be performed in accordance with several embodiments. aspect of this first embodiment, the building blocks are mononucleotides, however In a first embodiment the step of contacting the complementing element with the dance with this embodiment the building blocks is a nucleotide derivative. In one ဓ 33

well as methods for preparing such compositions and uses thereof, wherein each of said compositions comprise a templated molecule according to the Invention in an

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amount effective to achieve a desired effect.

secticidal composition, a bacterlocidal composition, and a fungicidal composition, as

In other aspects of the invention there are provided a pesticidal composition, an in-

agent, or a diagnostic agent, wherein said method comprises the step of screening a In still further aspects there is provided a method for identifying a pharmaceutical

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natural substrate for polymerases and transscriptses, oligonucleotides are incorporable in accordance with the method of WO 01/16366. The mono- or oligonucleotide derivative serves as the complementing element. One or more linker(s) is/are attached at one end to the mono- or oligonucleotide derivative and at the other end to a functional entity. Especially, in the case in which the complementing element is a mononucleotide derivative, it is preferred that the linker is attached so that the functional entity is projecting into the major groove of a double stranded helix to allow adjacent functional entities to form a linkage to each other.

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In a second embodiment of the invention, building blocks comprising an mono- or oligonucleotide as complementing element are chemically ligated together. Several methods for chemical ligation are know in the art, such as the 5'-phosphoimidazolid method (Visscher, J.; Schwartz, A. W. Journal of Molecular Evolution 1988, 28, 3-6. And Zhao, Y.; Thorson, J. S. J. Org. Chem. 1998, 63, 7568-7572) or the 3'-

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phosphothloate method (Alvarez et al. J. Org. Chem. (1999), 64, 6319-28 Pirrung et al. J. Org. Chem. (1998), 63, 241-46).

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In a third embodiment of the invention, building blocks comprising an oligonucleotide as complementing element is ligated together using a ligase enzyme.

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In a fourth embodiment of the invention, the building blocks comprise an oligonucleotide as complementing element, said oligonucleotide having a sufficient length to adhere to the template without the need for ligation to a primer or an other complementing element.

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The building blocks are in general adapted to the method used for contacting the complementing element with the template and production of the templated motecule. As an example, the linker may be relatively short when a mononucleotide derivative is used, while the linker needs to be considerable longer when an oligonucleotide is used as building block.

Brief Description of the Figures

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The following symbols are used in the following figures to indicate general characteristics of the system: In figures 1, 7C, 8C, 11, 11 ex.1, 12, 13, 14, 14 ex. 1 – 2, 15, 15 ex. 1 – 7, 17, 17 ex. 1, 17, 17 ex. 1 – 2, 19, 19 ex. 1 – 3, 20, 21, and 22A, a long horizontal line symbolizes a template, complementing template or the complex of the template with the complementing template. For clarity, in some of the figures only the polymerization step, not the activation step, has been included. Rx denotes

Figure 1. Chemical Display of Templated Molecules - The principle.

unctional groups.

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The protocol for the chemical display of templated molecules can be divided into 6 steps, i) incorporation, ii) polymerization, iii) activation, iv) selection/screening, v) amplification, and vi) characterization, incorporation involves the incorporation of building blocks into the complementing template, which sequence is determined by the template.

Incorporation may be mediated by enzymes such as polymerase or ligase. The template comprises primer binding sites at one or both ends (allowing the amplification of the template). The remaining portion of the template may be of random, partly random or predetermined sequence. The complementing elements preferably comprises of a functional entity, a complementing element and a linker connecting

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20 the functional entity and the complementing element. Detailed examples of selected complementing elements, their incorporation, polymerization and activation are shown in (Figure 7 and 8).

Polymerization involves reactions between the incorporated building blocks, thereby forming covalent bonds between the functional entities, in addition to the functional bonds that already exist between the complementing elements.

Activation involves cleaving some, all but one, or all of the linkers that connect the sequence of functional entities to the template or complementing template having templated the templated molecule comprising the functional entities. Activation may also involve separating the template and the complementing template without cleaving the linkers connecting the functional entities and the complementing

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template.
Selection or screening involves enriching the population of template-templated molecule pairs for a desired property.

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Cloning and sequencing involves the cloning of the isolated templates or complementing templates, followed by characterization. In some cases, it may be desirable to sequence the population of isolated templates or complementing templates, wherefore cloning of individual sequences are not required.

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10 Figure 2A and 2B. An expanded set of base pairs.

The figure discloses a set of natural and non-natural base pairs that obeys Watson-Crick hydrogen-bonding rules. The base pairs are disclosed in US 6,037,120, incorporated herein by reference.

15 Figure 3. A monomer building block.

A building block comprises or essentially consists of a functional entity, connected through a selectively cleavable linker to a complementing element. Each complementing element has two reactive groups (type I), which may react with two other complementing element contains a recognition group that interacts with a complementary coding element contains a recognition group that interacts with a complementary coding element contains of shown). The functional entity in this example comprises or essentially consists of two reactive groups (type II), which may react with reactive groups of other functional entitie(s), and a functional group, also called a functionality. The reactive groups of type II, and the molecular molety that connects them, will become (part of) the backbone in the resulting encoded polymer.

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Figure 4. A monomer building block with only one reactive group type II.

A building block comprises or essentially consists of a functional entity, connected through a selectively cleavable linker to a complementing element. Each complementing element has two reactive groups (type I), which may react with other complementing elements. The complementing element (coding element not shown).

The functional entity in this example comprises or essentially consists of a reactive group type II, which may react with reactive groups of other functional entities, and a

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functional group, also called a functionality. The reactive group type II will become (part of) the backbone in the resulting encoded polymer.

Figure 5. Building blocks and the polymers resulting from template directed incorporation of the building blocks and their polymerization and activation Figure 3 discloses a detailed description of features of individual building blocks.

Three different complementing elements are shown, each linked to a specific functional entity. The right half of the figure includes the template which directs the incorporation of the building blocks by complementary base paining.

10 A). The reactive groups type I of the complementing element react, whereby a part of the reactive group is lost (e.g., PPi in the incorporation of nucleoside triphosphates). In the shown example, the polymerization of reactive groups type II also results in loss of part of the reactive groups. The backbone of the resulting polymer comprises or essentially consists of part of the original reactive groups type II and the molecular entity that connects the reactive groups. Part of the linker remains attached to the functional entity.

B). The reactive groups type I react as in (A). The reactive groups type If do not react directly, but rather a "bridging molecule" is added. Upon reaction with this bridging molecule, part of the reactive group is lost. The cleavable linker used in this example is a so-called "traceless linker" and therefore the functional entity is

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released with no trace of the linker molecule.

C). Incorporation in this case does not involve coupling of the individual complementing elements, i.e., does not lead to the reaction of the reactive groups type I. The reactive groups type II react with bridging molecules as in (B).

25 D). The functional entity contains only one reactive group type II. The reactive group type II reacts with a bridging molecule.

Figure 6. A derivatized nucleotide as building block

The nucleotide building block comprises or essentially consists of the complementling element (the nucleotide) and a functional entity (in this case a dicarboxylic acid)
connected by means of a selectively cleavable linker (here a disulfide). The reactive
groups type I of the nucleotide are the triphosphate and the hydroxyl group, as indicated. The recognition group of the nucleotide is the base. The functional entity
comprises or essentially consists of a functional group (a hydroxyl), two reactive
groups type II (carboxylic acids), and a backbone structure (aromatic ring) connect-

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ing the two reactive groups. Finally the linker (disulfide) is cleavable by for example Ë

A derivatized di-nucleotide as building block

basic cleavage, the linker releases the functional group, a carboxylic acid. The reaclive groups of type I of the di-nucleotide are the hydroxyl group and the phophoro(2methyl)imidazolide. Reactive groups of type II are the amino group and the carboxfunctional entity (here an amino acid) via a cleavable propargylester linkage. Upon The complementing element is a modified dA-dU di-nucleotide that comprises the recognition group, in this case the adenine and uracil bases. It is connected to the /lic acid of the amino acid as indicated. ໝ

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A derivatized oilgo-nucleotide as building block

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hexane spacer connected to an N-hydroxysuccinimid molety. Reactive group of type linked to the functional entity, a M-Boc beta amino acld, via an oligo-nucleotide com-The complementing element is the last 20 bases of the oligonucleotide shown. It is able phosphoramidite (10-1953-95 from Glen research) including a cytosine deoxyphising 40 bases (B is an internal blotin incorporated using the commercially avail-Il is the carboxylic acld bound to the oxygen atom of the N-hydroxysuccinimid molribonucleotide that has been modified at the 5'-phosphato group with a mercaptoety. It is susceptible to nucleophillic attack by e.g. an amine.

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- Figure 7. C-terminal tagging of a β -dipeptide incorporation, polymerization and activation. 22
- The initiator is a Fmoc-protected amine. The dUTP-derivative carries a phothiocarboxyanhydride (NTA) ring structure. The dATP-derivative is modified A) Structures of the primer and two monomer building blocks. The initiator molecule is attached to the 5-position of the 3'-terminal dU of the primer. toprotected hydroxyl group. The hydroxyl group is coupled to the Nat the 7 position. A photoprotected amine is coupled the NTA.

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tended from its 3'-end through incorporation of the dUTP and dATP by a po-The primer (which is annealed to the template, not shown in figure) is exlymerase. Then the initiator is activated by piperidine, which releases the

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ray. As a result, a polymer, attached through its functional groups (OH and primary amine. This primary amine now attacks the next NTA unit in the aropens the NTA rings structure, releases CSO, and as a result, produces a primary amine. The primary amine attacks the neighboring NTA, which

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- strand with the NTA units, are cleaved. The resulting polymer in this case is Natures encoding system where 5' to 3' RNA encodes an $\alpha\text{-peptide}$ in the N-DNA sequence dUdA. In the shown example, the sequence 5'-dUdA-3' en-NH2) to the DNA strand, Is formed. Finally, the linkers connecting the DNA codes a β -peptide in the C-terminal to N-terminal direction, as opposed to a β-peptide, carrying the functional groups OH and NH2, encoded by the to C-terminal direction. The β-peptide is attached to the encoding DNA
- C) Schematic representation of the Incorporation, polymerisation and activation. The encoded polymer becomes attached to the encoding molecule (DNA) through the initiator molecule.

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through its C-terminal end.

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Figure 8. N-teminal tagging of a β-dipeptide - incorporation, polymerization and activation.

- The Initiator is a Fmoc-protected amine. The UTP-derivative carries a photothiocarboxyanhydride (NTA) ring structure. The ATP-derivative is modified A) Structures of the primer, two monomer building blocks, and an oligo. The primer. The primer is complementary to the upstream part of the template. initiator molecule is attached to the 5-position of the 3'-terminal U of the protected hydroxyl group. The hydroxyl group is attached to the N-22 2
- ty of the thioester in water can be modified as desired by changing the strucat the 7 position. A photo-protected amine is attached to the NTA. The oligo carries a reactive thioester attached to the U at the oligo's 5'end. The stabilture of the thioester-component (in the example, the thiol-component is a is complementary to the downstream sequence of the template. The oligo
- merase. Then the initiator is activated by piperidine, which releases the primary amine. The primary amine attacks the neighboring NTA, which opens tended from its 3'-end through incorporation of the UTP and ATP by a poly-B) The primer (which is annealed to the template, not shown in figure) is ex-

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the NTA rings structure, releases CSO, and as a result, produces a primary amine. This primary amine now attacks the next NTA unit in the array. As a result, a polymer, attached through its functional groups (OH and NH₂) to the RNA strand, is formed. Finally, the linkers connecting the RNA strand with the NTA units are cleaved. The resulting polymer is a β-peptide, carrying the functional groups -OH and -NH₂, encoded by the ribonucleic acid sequence UA. The sequence 5'-UA-3' encodes a β-dipeptide in the N-terminal to C-terminal direction, similar to the way that Nature encodes α-peptides. The β-peptide is attached to the encoding RNA through its N-terminal end.

C) Schematic representation of the incorporation, polymerisation and activation. Upon cleavage of a subset of linkers, the encoded polymer becomes attached to the downstream oligonucleotide.

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Figure 9. Nucleotide—derivatives that are known to be incorporated into RNA or DNA strands by DNA or RNA polymerases.

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Top: Nucleotide, the four bases and the site of attachment of the molecular moiety (R).

Center: Nucleotides with appendices (R) that are accepted as substrates by

polymerases.

Bottom: Nucleotides with appendices (R) that may be used with the present invention. Compound (a) would be used in for example fill-in experiments (see Figure 15). Compound (b) would be used for example in zipping polymerization reactions (see Figure 14 and 14, example 1). Compound (c) would be used for example in ring-opening polymerization reactions (see Figure 18 and 18, example

Figure 10. Cleavable Ilnkers and protection groups.

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Cleavable linkers and protection groups, agents that may be used for their cleavage and the products of cleavage.

Figure 11. Polymerization by reaction between neighboring reactive groups type II.

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For clarity, only the polymerization reaction (and not the activation) is shown in the figure. X represents the reactive groups type II of the functional entity. In this case the two reactive groups type II are identical.

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Polymerization (reaction of X with X to form XX) either happens spontaneously when the monomer building block has been incorporated, or is induced by a change of conditions (e.g. pH), or by the addition of an inducing factor (chemical or UV exposure, for example)

Figure 11 ex.1. Coumarin-based polymerization.

Light-induced reaction of the coumarin units, followed by activation (cleavage of the linker), results in a polymer backbone of aromatic and aliphatic ring structures. Examples of functional groups (phosphate, carboxylic acid and aniline) are shown.

Figure 12. Polymerization between neighboring non-identical reactive groups type II.

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In this example, X may react with Y but not another X. Likewise, Y does not react with Y. Polymerization can either happen during the incorporation of building blocks (as shown in the figure), or after incorporation of several building blocks.

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Figure 13. Cluster formation in the absence of directional polymerisation. When the incorporated monomers are not fixed with regard to rotation about the bond that links the functional entities to the complementing elements, cluster formation may result, as shown in the figure.

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This represents a significant problem for longer polymers. The problem may be solved by (i) fixing the incorporated monomers in a preferred orientation which does not allow X and Y (reactive groups type II) to exchange positions in the array (e.g., by coupling the functional entity and the complementing element via a double bond or two honds are counting the functional entity and the complements.

or two bonds, e.g.. coupling the functional entity both to the base and the ribose of a nucleotide, or to the two bases of a dinucleotide), (ii) employing directional polymerisation ("zipping", see for example figure 17), or (iii) setting up conditions that ensure that the monomers react during or right after incorporation into the complementing template, i.e., each monomer reacts with the previously incorporated

30 monomer before the next monomer is incorporated (see for example Figure 14, with example).

Figure 14. Zipping-polymerization and simultaneous activation.

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between X and Y is in this example the same for all monomers participating in the Polymerization results In activation of the polymer. The geometry of the reaction

Figure 14, example 1. Simultaneous Incorporation, polymerisation and activation - formation of peptides.

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the next monomer is incorporated, this may attack the thioester carbonyl, resulting in cleavage of the dipeptide from the nucleotide, to form a tripeptide. The process conare Incorporated. During or after incorporation of a nucleotide-derivative, the amine attacks the carbonyl of the (previously incorporated) neighboring nucleotide. This (A). Nucleotide derivatives, to which amino acids thioesters have been appended, results in formation of an amide bond, which extends the peptide one unit. When cleotide derivatives stops. Importantly, the geometry of the nucleophilic attack retinues further downstream the complementing template, until incorporation of nu-

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mains unchanged. As the iocal concentration of nucleophilic amines is much higher on the template than in solution, reactions in solution is not expected to significantly the reactivity include: (i) pH and temperature, (ii) length, point of attachment to the pho-, or hydroxy-ester); (iv) the nature of the substituent on the sulfur (see (B) below). In addition, the efficiency of correct polymer formation is also affected by the affect the formation of the correct encoded polymer. Furthermore, the reactivity of the amine with the ester may be tuned in several ways. Parameters that will affect rate of incorporation and rate of reaction once incorporated. The rate of incorporachanging the conditions (pH, concentration of nucleotides, salts, templates and enzymes), by choice of enzyme, or by changing the characteristics of the enzyme by protein engineering. Also, the nature and size of the nucleotide-derivatives may linker that connects the ester and the nucleotide, (iii) nature of ester (thio-, phosnucleotide, and characteristics (charge, rigidity, hydrophobicity, structure) of the tion is determined by kcat and Km. The kcat and Km values may be tuned by

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This general scheme involving incorporation, polymerisation and activation during or right after building block incorporation, can be applied to most nucleophilic polymensation reactions, Including formation of various types of peptides, amides, and amide-like polymers (e.g., mono-,di-, tri-, and tetra-substituted α -, β -, γ -, and Ω -

influence its rate of incorporation.

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peptides, polyesters, polycarbonate, polycarbarmate, polyurea), using similar struc-

(B). Four different thioesters with different substituents and therefore different reaciivity towards nucleophiles.

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Figure 14, example 2. Simultaneous incorporation, polymerization and activation - formation of a polyamine.

containing the nucleophilic center attacks the electrophile attached to the DNA-part, This figure shows a "rolling-circle polymerization reaction" where the chain

using this DNA-part as the leaving group. 9

Fill-in polymerization by reaction between reactive groups type II ("X" in the figure) Figure 15. "Fill-in" polymerization (symmetric XX monomers).

and bridging molecules (Y-Y) in figure).

analog. X represents the reactive groups type II of the functional entity. In this case ure. The thick line represents double or single stranded nucleic acid or nucleic acid during or after incorporation of the monomer building blocks. Likewise, significant For clarity, only the polymentzation reaction (not the activation) is shown in the figthe two reactive groups type II are identical. (Y-Y) is added to the mixture before, 5

reaction between X and Y may take place during or after incorporation of the nonomers. ឧ

Figure 15, ex.1. Poly-Imine formation by fill-in polymerization.

Dialdehyde is added in excess to incorporated diamines. As a result, a poly-imine is formed. In the example, the polymer carries the following sequence of functional groups: cyclopentadienyl, hydroxyl, and carboxylic acid.

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Figure 15, example 2. Polyamide formation.

tions. Atternatively, a di-(N-hydroxy-succinimide ester) may be added in excess, at a After incorporation of nucleotides to which have been appended di-amines, EDC (1pH of 7-10. As a result, two amide-bonds are formed between two neighboring nu-Ethyl-3-(3-dimethylaminopropyl)carbodiimide) and dicarboxylic acid is added in excess to the primary amines on the oligonucleotide using standard coupling condi-ဓ

cleotide-appendices. After this polymerisation, the appendices are separated from the oligonucleotide backbone (activation), leaving one linker intact, and the pro-35

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tected functional groups are deprotected to expose the functional groups. The final result is a DNA-tagged polyamide.

An alternative route to polyamides would be to incorporate nucleotides to which had been appended di-carboxylic acids, and then add di-amines and EDC, to form amide bonds between individual nucleotides of the oligonucleotide. Alternatively, the nucleotide derivatives might contain N-hydroxy-succinimidyl (NHS) esters, which would react with the added amines without the need to add EDC. Initially, this latter method was considered to be problematic in the case where incorporation is mediated by a polymerase, as the NHS-esters probably would react with amines on the polymerase, potentially inhibiting the activity of the polymerase. However, practical experiments have shown that it is possible to incorporate NHS-derivatised nucleotides.

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(A). The backbone of the resulting polymer comprises or essentially consists of amide-bonded aromatic rings. The substituents of this example are a protected primary amine, a branched pentyl group, a tertiary amine and a pyrimidyl. The primary amine is protected in order to avoid its reaction with the dicarboxylic acid. Appropriate protecting groups would be for example Boc-, Fmoc, benzyloxycarbonyl (Z, cbz), trifluoracetyl, phthaloyl, or other amino protecting groups described e.g. in (T. W. Green and Peter G. M. Wuts (1991), Protective Groups in Organic Synthesis).

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(B). The backbone comprises or essentially consists of aromatic rings, connected by amide bonds. The substituents are indanyl, diphenylphosphinyl, carboxamidoethyl and guanidypropyl, the latter two representing the asparagine side chain, and the arginine side chain, respectively. The guanidyl function is protected, as it is more reactive than standard amines. An appropriate protecting group would be Mit (4-methoxy-2,3,6-trimethylbenzenesulfonyl), Mits (mesitylene-2-sulfonyl) or Pbf (2,2,4,6,7-pentamethyldihydro-benzifuran-5-sulfonyl).

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Figure 15, example 3. Polyurea formation.

The Incorporated nucleotide derivatives react with phosgen or a phosgen-equivalent such as CDI to form a polyurea. The linkers are cleaved and the protected hydroxyl is deprotected.

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Appropriate leaving groups (Lv) are chloride, Imidazole, nitrotriazole, or other good leaving groups commonly employed in organic synthesis

35 Figure 15, example 4. Chiral and achiral polyurea backbone formation.

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In this example, the functional group Rx is used as a cleavable linker, that generates the desired functional group upon activation. In both (A) and (B), a polyurea is

In (A), the functional group is attached to the backbone via a chiral carbon. The hydrogen on this carbon is drawn to emphasize this. Before polymerisation, there is free rotation about the bond connecting the chiral carbon and the functional group. When the reactive groups type II (the amines) react with the phosgen equivalent (e.g., a carbonyidiimidazole) to form the polymer, the building blocks may be inserted in either of two orientations (as indicated by the position of the hydrogen, left

or right). As a result, each residue of the polymer has two possible chiral forms.

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Therefore, a given encoding molecule will encode a polymer with a specific sequence of residues, but an encoded polymer of 5 or 15 residues will have $2^5 = 32$ or $2^{15} = 32768$ stereoisomers, respectively. In certain cases it may be advantageous to incorporate such additional structural diversity in the library (for example when the polymer is relatively short). In other cases such additional diversity is not desirable, as the screening efficiency may become compromised, or it may become too difficult to deconvolute the structure of a polymer that has been isolated in a screening process, together with the other stereolsomers encoded by the same encoding

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In (B), the chiral carbon of (A) has been replaced by a nitrogen. As a result, the resulting polymer backbone is achiral, and the encoding molecule encodes one specific structure.

molecule (for example when the polymer is long).

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Figure 15, exampte 5. Polyphosphodiester formation.

The incorporated nucleotide derivatives react with the activated phosphodiester to form a polyphosphodiester. Then the linkers are cleaved, resulting in a polyphosphodiester, attached through a linker to the encoding molecule.

An example of an appropriate leaving groups (Lv) is imidazole.

30 Figure 15, example 6. Polyphosphodlester formation with one reactive group type II in each monomer building block. Each incorporated nucleotide contains an activated phosphodiester. Upon addition of a dihydroxylated compound such as 1,3-dihydroxypyridine, a functionalised polyphosphodiester is formed. Finally, the functional groups Rx are liberated from the

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complementing template by cleavage of the protection groups/cleavable linker that connected them to the oligonucleotide.

Figure 15, example 7. Pericyciic, "fill-in" polymerization.

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structure is activated by cleaving the linkers that connect the polymer to the nucleo-After incorporation of the nucleotide-derivatives, 1,4-benzoquinone is added in excess, resulting in the formation of a polycyclic compound. Finally, the polymeric tides, except for one (non-cleavable) linker which is left intact.

Figure 16. Encoded "Fill-In". 9

is the Y-R_x-Y of the second building block. Using this method it is possible to link two Fill-in by encoding is performed by the method depicted. The encoded fill-in molety bodiments this may be of advantage because the encoded fill-in functional entity Y-Rx-Y does not have to be the same through out the molecule, as is the case for the functional entities X-R_x-X by a predetermined functional entity Y-R_x-Y. In some emmethod shown in Fig. 15.

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Figure 17. "Fill-In" polymerization (asymmetric XS monomers).

Fill-in polymerization by reaction between reactive groups type II ("X" and "S" in the figure) and bridging molecules (T-Y) in figure).

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ing or after incorporation of the monomer building blocks. Likewise, significant reacline represents double or single stranded nucleic acid or nucleic acid analog. X and tion between X and Y, and between S and T may take place during or after incorpo-S represent the reactive groups type II of the functional entity. In this case the two For clarity, only the polymerization reaction (not the activation) is shown. The thick reactive groups type II are non-identical. (T-Y) is added to the mixture before, durration of the monomers.

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Figure 17, example 1. Fill-in polymerization by modified Staudinger ligation

and ketone-hydrazide reaction. ဓ

The reactive groups (type II) X and S of the building blocks are azide and hydrazide between a azide and a phosphine, are very chemoselective. Therefore, most functional groups Rx can be employed without the need for protection during the polym-The added molecule that fills the gaps between the building blocks carry a ketone and a phosphine mojety. The reactions between a ketone and a hydrazide, and

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found in (Mahal et al. (1997), Science 276, pp. 1125-1128; Saxon et al. (2000), Orerization reactions. Examples for the molecular moieties R, R1, X and Y may be ganic Letters 2, pp. 2141-2143).

Figure 18. "Zipping" polymerization. 2

The initiator molecule (typically located at one of the ends of the nascent polymer) is active group Y for attack on the neighbouring X. Polymerisation then travels to the acts with the reactive group X of the neighbouring monomer. This activates the reactivated, for example by deprotection or by a change in pH. The initiator then re-

other end of the molecule in a "zipping" fashion, until all the desired monomers have for attack by it on the neighbouring reactive group, or activation of it for attack by the been connected. The activation of the initiator (and reactive groups Y) may be both neighbouring reactive group. 9

Figure 18, example 1. Radical polymerisation. 5 The initiator molecule, an iodide, is activated by the addition of a radical initiator, for example ammonium persulfate, AIBN (azobis-isobutyronitrile) or other radical chain radical and a bond between the first two monomers. Eventually the whole polymer reaction initiators. The radical attacks the neighboring monomer, to form a new

is formed, and the polymer may be activated, which simultaneously creates the unctional groups Rx. ឧ

Figure 18, example 2. Catlonic polymerisation.

A cation is created by the exposure of the array to strong Lewis acid. The double charge migrates to the neighbouring monomer. Eventually the whole polymer is bond of the neighbouring monomer reacts with this cation, whereby the positive formed, and finally it is activated.

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Figure 19. Zippting polymerization by ring opening.

ring, whereby the reactive group Y in the same functional entity is activated for reac-The initiator reacts with the reactive group X in the ring structure, which opens the ion with a reactive group X in a neighboring functional entity. ဓ္တ

Figure 19, example 1. "Zipping" polymerization of N-thiocarboxyanhydrides,

to form β-peptides. 33

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A number of changes to this set-up can be envisaged. For example, instead of thioprotected with a base- or photolabile group. If a base-labile protection group is cho-Finally, the initiator might be unprotected and for example coupled to the primer. In amine then attacks the carbonyl of the neighbouring N-thiocarboxyanhydride (NTA) blocks the local concentration of initiator and carboxyanhydride will be much higher. sen, the stability of the carboxyanhydride must be considered. At higher pH It may unit. As a result, CSO is released, and a primary amine is generated. This amine will now react with the next NTA unit in the array, and eventually all the NTA units After incorporation of the building blocks, the initiator is deprotected. The primary carboxyanhydrides, one might use carboboxyanhydrides. The initiator might be nanomolar to micromolar), wherefore only an insignificant amount of initiator will be advantageous to use carboxyanhydrides rather than thiocarboxyanhydrides. react with the carboxyanhydrides. After or during incorporation of the building this case the concentration of the initiator in solution will be very low (typically will have reacted, to form a b-peptide. Finally, the oligomer is activated. leading to efficient polymerization.

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polyurea) can be made, using similar cyclic structures. For example, α-peptides can Other types of peptides and peptide-like polymers (e.g., mono-,di-, trl-, and tetrasubstituted α , β , γ , and Ω -peptides, polyesters, polycarbonate, polycarbamate, be made by polymerization of 5-membered carboxyanhydride rings.

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Figure 19, example 2. "Zipping" polymerization of 2,2-diphenylthiazinanone units to form p-paptides.

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The deprotected nucleophile, a primary amine, attacks the carbonyl of the neighborform a thioketone. As a result a free primary amine is generated, which attacks the around the secondary amine, making it more nucleophilic, whereas the nucleophilic ample, replacing the two aromatic rings with one aromatic ring will decrese the bulk formed, linked through its C-terminal end. The reactivity of the primary amine with ester), pH during the polymerization reaction and the choice of substituents on the by the bulk at the carbon between the secondary amine and the thioester. For excyclic molety and the primary amine released upon ring-opening, may be adjusted ing thioester, thereby forming an amide bond. The released thiol reorganizes, to the ester may be modified for example by the choice of ester (thicester or regular carbonyl of a neighboring thioester, etc. Eventually an α -substituted β -peptide is aromatic ring(s). The relative reactivity of the secondary amine contained in the

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principle. For example, 1-peptides may be formed by polymerization of 7-membered ity of the primary amine that is formed upon ring-opening is not affected by the bulk at this position. Other peptides and amide-like polymers may be formed by this thlazinanone rings.

Figure 19, example 3. Polyether formation by ring-opening polymertsation.

The initiator is deprotected by for example base or acid. The formed anion the nattacks the epoxide of the neighboring monomer, to form a ether-bond. As a result,

conditions the attack will be at the most or least hindered carbon of the epoxide (unarray, and eventually the full-length polyether has been formed. Depending on the an anion is formed in the neighboring unit. This attacks the next monomer in the der acidic or basic conditions, respectively). 9

in the final step, the encoded polyether is activated. In this case, the polymer is fully released from the encoding molecule. The screening for relevant characteristics

and therefore the templates encoding polyethers with interesting characteristics may (e.g., effect in a cell-based assay or enzymatic activity) may be performed in microiter wells or micelles, each compartment containing a specific template molecule plated molecule is physically associated (by the boundaries of the compartment), and the templated polyether, in many copies. In this way, the template and tem-5

be collected from those compartments, pooled, amplified and "translated" Into more copies of polyethers which may then be exposed to a new round of screening. ଷ୍

Figure 20. Zipping-polymerization and activation by rearrangement.

The initiator is activated for attack by Y. Reaction of initiator and Y results in release rearrangement of the building block molecule takes place, resulting in activation of X for reaction with Y. After a number of reactions and rearrangements, a polymer has of the initiator from the complementing element. Upon reaction with the Initiator, a been formed. 23

Reaction of the initiator with X in the ring structure opens the ring, resulting in activaring-opening, the functional entities are released from the complementing elements. ion of Y. Y can now react with X in a neighboring functional entity. As a result of Figure 21. Zipping-polymerization and activation by ring opening. ഉ

Figure 22. Directional polymer formation using fixed functional units.

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around the linker that connects functional entity and complementing element (A) The functional entity of a building block may be attached to the complementing element through two linkers. This may fix the functional entity in a given (as depicted in figure 13) Is not possible, and cluster formation therefore orientation relative to the complementing template. As a result, rotation

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(B) Two linkers connect the two bases of a dinucleotide-derivative with the func-(A) above) In proximity to the ester (Y In (A) above). This ester may be actithe polymerisation reaction will cleave the ester from the base to which it is of the linker that connects the base at the 3'-end of the dinucleotide with the amine and the ester, a polypeptide is formed. This polypeptide will be a ditional unit, which in this case is a dipeptide. Incorporation of such dinucleotide derivatives into a double helical structure will position the amine (X in vated, for example as a N-hydroxysuccinimide ester. After reaction of the rectional polymer, with N-to-C-terminal directionality. In the present case, inked. Therefore, activation of the formed polymer only requires cleavage amino-terminal end of the functional entity.

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may be coupled to the complementing element through a double bond, or it Rotational fixation of the functional entity relative to the complementing elebose or base. Finally, it is also possible to link to the phosphate moity, esmay be attached through two bonds to the base and ribose molty of a nucleotide, respectively, or it may be coupled to different positions on the riment may be achleved in other ways. For example, the functional entity pecially of a dinucleotide.

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(A), on the sugar moiety of the parent nucleotide (B, linked through ester functionalty or C, with ester functionality free, and D, also with ester functionality free), it can Figure 23 shows four examples of bifunctional FEs attached via a single linker to point (right). The second attachment can be anywhere on a neighbour nucleotide be another base position of the parent nucleotide (not shown), or the FE could be the parent nucleotide (left) or with an additional linker using a second attachment linked to the phosphate backbone (not shown).

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linker-FE 1A attached in different ways. DNA backbones are shown as arrows, sugcoloured by atom. A. Example of a conformation bearing singly-attached FEs. B. ars and bases as rings. Linker-FE atoms are depicted in stick representation and Figure 24 show a DNA double helix (upper strand 5'-GCTTTTTAG-3') bearing

Most probable product of A. C. Example of a singly-attached FE configuration leading to clustering and thereby to an incomplete product, D. E. Minimum energy conformation bearing doubly-attached FEs and the only possible product, F. G. Stick representation of the released product from F. H. Stick representation of the reeased product from B. I. Stick representation of the released product from D. S

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FE 1B attached in different ways. DNA backbones are shown as arrows, sugars and Figure 25 show a DNA double helix (upper strand 5'-GCTTTTAG-3') bearing linkerbases as rings. Linker-FE atoms are depicted in stick representation and coloured

probable product of A. C. Example of a singly-attached FE configuration leading to clustering and thereby to an incomplete product, D. E. Minimum energy conformation bearing doubly-attached FEs and the only possible product, F. G. Stick repreby atom. A. Minimum energy conformation bearing sIngly-attached FEs. B. Most sentation of the released product from B and F. H. Stick representation of the reeased product from D. 5

Figure 26. Templating of molecules - principle and variations.

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In the figures 26-27, 29-31, 33-35, 37-49, and 53, the template, the complementing template, both the template and the complementing template, or a complementing element is Indicated by a horizontal (bold) line. In figures 26-28, 35-37, and 39, a circle is used to indicate a functional entity.

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A. Monomer bullding blocks used in this figure. A black dot indicates a cleavable linker.

General principle. œ.

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porated into a complimentary template, by specific Interaction between cod-Step 1 - Incorporation. The monomer building blocks are specifically incoring elements (of the template) and complementing elements (of the mono-

mer building blocks).

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Step 2 – Reaction. A reaction is induced by which functional entities (FE) of the individual monomer bullding blocks become coupled, by reaction of reactive groups type II.

Step 3 – Activation. Some or all of the linkers connecting the FE units with complementing elements are cleaved, thereby partly or fully releasing the templated motecule.

Step 4 (not shown In figure) – Screening, Amplification and Modification.

The template-templated molecule complexes may be taken through a screening process that enriches the pool for complexes with desired features. Then the templates of the enriched pool may be amplified and modified, by e.g. mutagenic PCR, and the templated molecules regenerated by performing step 1-3.

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C. Templating of linear, branched and circular templates.

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Linear, branched and circular templates may generate linear, branched and circular templated motecules. In the example shown, the branched template may be generated by incorporation of a modified nucleotide (e.g., carrying a thiol) into an oligonucleotide, followed by reaction with an oligonucleotide containing a thiol-reactive component (e.g., a maleimide-unit at one end). The circular template may likewise be a oligonucleotide, carrying reactive groups at the end that may react to covalently close the circle (e.g., thiols at both ends of the oligonucleotide could form an disulfide bond). Upon cleavage of all but one of the linkers connecting the FEs and complementing elements, a circular templated molecule is formed, attached to the template at one point.

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 D. Templating of linear, branched, circular and scrambled linear molecules by linear template.

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(a) A linear templated molecule with the same sequence of FEs as obtained after incorporation, but before reaction, of the monomer building blocks. (b) A linear templated molecule with a scrambled sequence, i.e., the sequence of the FEs in the templated molecule does not correspond to the sequence obtained right after incorporation, but before reaction of the FEs. (c) A circular templated molecule obtained by pairwise reaction of the following FEs with each other: FE1/FE2, FE2/FE3, FE3/FE5, FE5/FE4, FE4/FE1. (d) A branched molecule obtained by pairwise reaction of the following functional

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entities with each other. FE1/FE2, FE2/FE3, FE2/FE4, and FE4/FE5. (e) A branched molecule obtained by paliwise reaction of the following functional entities with each other. FE1/FE2, FE2/FE4, FE2/FE5, FE2/FE3.

Figure 27. Non-equal number of reactive groups (X) and (Y). The number of reactive groups (X) can be higher than, equal to, or lower than the number of reactive groups (Y). When the number of (X) and (Y) are different, scrambling results. In the figure the scaffold (the molecular moiety to which the functional groups of the monomer building blocks become attached) is directly attached to the template. The scaffold may also be part of a monomer building block (i.e., the functional entity of the monomer building block comprises a scaffold molety, including reactive groups

(A). Number of encoded reactive groups X per template equals the number of reactive groups (Y) on the anchorage point (also called the scaffold).

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(B). Number of encoded reactive groups X per template is less than the number of encodable substitutent positions Y on the scaffold. This leads to scrambling regarding which of the reactive groups (Y) on the scaffold (anchorage point) will react with an (X) on the monomer building blocks.

C. Number of encoded reactive groups X per template is larger than the number of reactive groups on the scaffold. This leads to scrambling regarding which of the reactive groups (Y) on the scaffold (anchorage point) will react with a reactive group (X) on the monomer building blocks.

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Figure 28. Monomer building blocks.

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(A) A monomer building block with one reactive group type II (X), connecting the functional group (Rx) with the complementing element. This type of monomer building block may be used for the simultaneous reaction and activation protocol (Figure 14).

(B) A monomer building block with two reactive groups type II (X and Y), connecting the complementing element and the functional group (Rx).

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(C) A monomer building block with one reactive group type II (X). The reactive group (X) does not link the functional group (Rx) and the complementing element, wherefore a linker (L) is needed for the activation step (in order to release the functional entity from the complementing element)

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(D) A monomer building block with four reactive groups type II (Y). The four reactive groups and the functional group Rx may serve as a scaffold, onto which substituents (encoded by monomers complementinig the same template) are coupled through reaction of reactive groups (X) on these monomer building blocks with the reactive groups (Y) on this monomer building block. In this example, no cleavable linker is indicated. Therefore, after the templating reactions the templated molecule may be attached to the template through the linker of this monomer building block.

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Figure 29. Templating involving simultaneous reaction and activation.

Templating using 4 monomer building blocks each with one reactive group type II

(X), and an anchorage point carrying 4 reactive groups (Y). The reaction of X and Y involves simultaneous activation (cleavage) which releases X from the complementing element.

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(A) The reactive groups type II (X) are of similar kind.

(B) The reactive groups type II (X1, X2, X3, X4) are of different kinds, i.e. the pairwise reactions between reactions X1/Y1, X2/Y2, X3/Y3, and X4/Y4 are orthogonal or partly orthogonal. For example, X1 preferably reacts with Y1, not Y2, Y3 or Y4. The anchorage point may be attached directly to the template, or to the complementing template. In case the anchorage point is attached to a complementing element, as a whole it is considered a monomer building block.

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Figure 30. Reaction types allowing simultaneous reaction and activation. Different classes of reactions are shown which mediate translocation of a functional group from one monomer building block to another, or to an anchorage point. The reactions have been grouped into three different classes: Nucleophilic substitutions, addition-elimination reactions, and transition metal catalyzed reactions. These reactions are compatible with simultaneous reaction and activation (as described in gen-

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(A) Reaction of nucleophiles with carbonyls. As a result of the nucleophilic substitution, the functional group R is translocated to the monomer building block initially carrying the nucleophile.

eral terms in figure 14).

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(B) Nucleophilic attack by the amine on the thioester leads to formation of an amide bond, in effect translocating the functional group R of the thioester to the other monomer building block.

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(C) Reaction between hydrazine and β-ketoester leads to formation of pyrazolone, in effect translocating the R and R' functional groups to the other monomer building block.

(D) Reaction of hydroxylamine with β-ketoester leads to formation of the isoxazolone, thereby translocating the R and R' groups to the other monomer building block. (E) Reaction of thiourea with β-ketoester leads to formation of the pyrimidine, thereby translocating the R and R' groups to the other monomer building

(F) Reaction of urea with melonate leads to formation of pyrimidine, thereby translocating the R group to the other monomer building block.

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(G) Depending on whether Z = O or Z = NH, a Heck reaction followed by a nucleophilic substitution leads to formation of cournarin or quinolinon, thereby translocating the R and R' groups to the other monomer building block.

(H) Reaction of hydrazine and phthalimides leads to formation of phthalhydrazide, thereby translocating the R and R' groups to the other monomer building block.

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 (i) Reaction of amino acid esters leads to formation of diketopiperazine, thereby translocating the R group to the other monomer building block. (J) Reaction of urea with a-substituted esters leads to formation of hydantoin, and translocation of the R and R' groups to the other monomer building block.

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(K) Alkylation may be achieved by reaction of various nucleophiles with sulfonates. This translocates the functional groups R and R' to the other monomer building block.

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leaving group, whereby the alkene is translocated to the nucleophile. (M) Reaction of disulfide with mercaptane leads to formation of a disulfide,

(L) Reaction of a di-activated alkene containing an electron withdrawing and a

(M) Neaction of disultide with mercaptane leads to formation of a disulfide, thereby translocating the R' group to the other monomer building block.

(N) Reaction of amino acid esters and amino ketones leads to formation of banzodiazepinone, thereby translocating the R group to the other monomer building block.

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(O) Reaction of phosphonates with aldehydes or ketones leads to formation of substituted alkenes, thereby translocating the R" group to the other monomer building block.

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- (P) Reaction of boronates with aryls or heteroaryls results in transfer of an aryl group to the other monomer building block (to form a biaryl).
- (Q) Reaction arylsulfonates with boronates leads to transfer of the aryl group.
- (R) Reaction of boronates with vinyls (or alkynes) results in transfer of an aryl group to the other monomer building block to form a vinylarene (or al-

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- (S) Reaction between aliphatic boronates and arylhalides, whereby the alkyl group is translocated to yield an alkylarene.
- (T) Transition metal catalysed alpha-alkylation through reaction between an enolether and an arylhallide, thereby translocating the aliphatic part.

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- (U) Condensations between e.g. enamines or enclethers with aldehydes leading to formation of alpha-hydroxy carbonyls or alpha, beta-unsaturated carbonyls. The reaction translocates the nucleophilic part.
- (V) Alkylation of alkylhalides by e.g. enamines or enolethers. The reaction translocates the nucleophilic part.
- (W) [2+4] cycloadditions, translocating the diene-part.

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- (X) [2+4] cycloadditions, translocating the ene-part.
- (Y) [3+2] cycloadditions between azides and alkenes, leading to triazoles by translocation of the ene-part.
- (Z) [3+2] cycloadditions between nitriloxides and alkenes, leading to isoxazoles by translocation of the ene-part. 8

Reaction of reactive groups (type II), followed by cleavage of the linkers that Figure 31. Templating involving non-simultaneous reaction and activation: connect functional entities with complementing elements. 22

(X), and an anchorage point carrying 4 reactive groups (Y). The reaction of X and Y and Y is followed by cleavage of the linker L, which releases the functional group Rx Templating using 4 monomer building blocks each with one reactive group type II does not involve simultaneous activation (cleavage), wherefore the reaction of X from the complementing element.

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are orthogonal or partly orthogonal. For example, X1 preferably reacts with Y1, not Y2, Y3 or Y4. The anchorage point may be attached directly to the template, or to (A) The reactive groups type II (X) are of similar kind, i.e., they may react with the same type of reactive group (Y). (B) The reactive groups type II (X1, X2, X3, X4) are of different kinds, i.e. the reactions between X1/Y1, X2/Y2, X3/Y3, and X4/Y4

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the complementing template. In case the anchorage point is attached to a complementing element, as a whole it is considered a monomer building block.

Figure 32. Pairs of reactive groups (X) and (Y), and the resulting bond (XY).

A collection of reactive groups that may be used for templated synthesis are shown, along with the bonds formed upon their reaction. After reaction, activation (cleavage) may be required (see Figure 31).

Figure 33. Anchorage sites for the templated molecule.

The templated molecule may be attached to the template that encodes it (A) through a linker that is connected directly to the template near the end of the template, or (B) through a linker that is connected directly to the template, at a more central position on the template, or (C) by way of a monomer building block carrying the anchorage point (a reactive group that becomes the linkage to the templated molecule). 9

Figure 34. Scrambling.

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blocks, the position or sequence of functional groups in the templated molecule may When the functional entities react after incorporation of the monomer building not always be uniquely determined by the template sequence. (1) The functional groups R1, R2, R3, and R4 may take any of the four positions on the scaffold molecule (i.e., the reactive group X of a monomer building block may react with any of the reactive groups Y on the anchorage point.

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(2) The sequence of one arm of this branched molecule may be e.g. R5-R3-R2 other of a number of possible sequences. Also, the Identity of the functional group coupled to e.g. the left part of the molecule, may be either of any of (as shown), or R5-R2-R3 (not shown), or R5-R4-R3 (not shown), or any

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- R1, R2, R3, or R4.
- (3) As in (2), a number of possible sequences of functional groups are possible, (4) Here a non-scrambled templated molecule is shown, in which the sequence in addition to the shown sequence R1-R2-R5-R4-R3.

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of the functional entitities when incorporated corresponds to the sequence of the templated molecule (R1-R2-R3-R4-R5). When desired, scrambling may be partly or fully avoided by directional encoding or the use of for example zipper boxes in the linkers (see figures 40, 44-47).

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(5) As in (2) and (3), a number of possible sequences and positions of the functional entities are possible.

Figure 35. Monomer building blocks - examples of linker design.

Different designs of monomer building blocks are shown, used in various schemes of termination.

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The complementing element may be represented by an oligonucleotide, to which a linker carrying the functional entity is attached. The linker may occupy an internal position with respect to the complementing element or alternatively occupy a terminal position. Both the complementing element and the linker may be made up of an oligonucleotide (DNA, RNA, LNA, PNA, other oligomers capable of hybridizing to the linker of a monormar building block and mixtures thereof). The horizontal part represents the complementing element, and the vertical part represents the linker.

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The portion of the linker marked "a" may be present or absent. Region "a" represents an interaction region of which one preferred embodiment is a sequence of nucleotides. Region "a" may be annealed to a complementary single stranded nucleotides sequence "a" in order to make the linker more rigid. Alternatively region "a" may be used for interaction with other monomer building blocks (i.e. zipper box see fig. 42), whereby the functional entities of such two monomer building blocks will be brought in close proximity, which will increase probability of reaction between these two functional entities. Other uses of such regions includes interaction between different monomer building blocks whereby directional encoding may be achieved. "Nu" is a nucleophile that may react with an electrophile "E".

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Different designs of monomer building blocks are shown, used in various schemes

25 of templating.

(A) The complementing element may be an oligonucleotide, to which a linker carrying the functional entity is attached to the central part of the oligonucleotide. The portion of the linker marked "a" may represent a nucleotide sequence to which a single stranded nucleotide may be annealed in order to make the linker more rigid.

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(B) Both the complementing element and the linker may be made up of an oil-gonucleotide. The horizontal part here represents the complementing element, and the vertical part represents the linker. The linker may contain a sequence "a" that functions as a zipper box (see figure 42).

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(C) The monomer building blocks of (C) is an initiator or anchorage point which may be used to initiate the encoding monoes.

may be used to initiate the encoding process.

Figure 36 Preparation of functional entities to oligonucleotide-based mono-

The reactions and reagents are shown that may be used for the coupling of functional entities to modified oligonucleotides (modified with thiol, carboxylic acid, halide, or amine), without significant reaction with the unmodified part of the oligonucleotide or alternatively, connective reactions for linkage of linkers to complementing elements.

Commercially, mononucleotides are available for the production of starting oligonu-

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deotides with the modifications mentloned.

Figure 37 Ollgonucteottde-based monomer building blocks. Examples of linker and functional entity (FE) design and synthesis.

- Examples are shown where the complementing elements of the monomer building blocks comparises oligonucleotides of length e.g. 8-20 nucleotides (oligonucleotide is drawn as a thick black line). Part of or all of the oligonucleotide may comprise the complementing element. In the case where only part of the oligonucleotide represents the complementing element, the remaining portion of the oligonucleotide may constitute a linker. In the examples, a linker is attached to the base on the 3'- or 5'- end of the oligonucleotide. This linker may be attached on any nucleotide in the
- constitute a linker. In the examples, a linker is attached to the base on the 3'- or 5'end of the oligonucleotide. This linker may be attached on any nucleotide in the
 oligonucleotide sequence, and also, it may be attached to any molecular molety on
 the oligonucleotide, as long as it does not abolish specific interaction of the complementing element with the template.
- (A) A monomer building block in which the linker (L) connects the base of the terminal nucleotide with the functional entity.

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- (B) A monomer building block in which a polyethylene glycol (PEG) linker of between one and twenty ethylene glycol units connects the complementing element with the functional entity which contains a nucleophile (a primary amine).
- (C) A monomer building block in which a linker (L) connects the functional entity which contains an electrophile (an ester or thioester).

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(D) A monomer building block comprising a Boc-protected amine (which may be deprotected with mild acid), and an ester. The deprotected amine may react with an ester of another monomer building block, to give an amide bond.

almost identical annealing temperatures wherefore mls-annealing is insignificant. In Figure 38. Oligonucleotide-based monomer building blocks. Example of coding and complementing element design, allowing for high monomer diversity. dom sequence (X specifies either C or G), and a constant sequence that is identical sequences (e.g., the sequences belonging to the group of BOX 1 sequences), have (A) Templaté carrying 6 coding elements (BOX 1-6), each containing a partly ranfor all sequences in the group (e.g., all BOX 1 sequences carry a central ATATTT sequence). By using C and G only (or, alternatively, A and T only), the individual

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groups of type II). The attachment point of the linker that connects the complementthe example, BOX 2 and BOX 3 are identical wherefore BOX 2 and BOX 3 may ening element and the functional entity is not specified in the figure. Ideally, the linker is attached to a nucleotide in the constant region, in order to avoid bias in the ancode the same type of functional entitles (comprising the same type of reactive

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(B) Example of coding element sequences. Example BOX 1 and BOX 6 sequences are shown. The example BOX1 sequence represents one specific sequence out of 1024 different sequences that anneal specifically to the corresponding BOX 1 complementing elements; the example BOX 6 sequence represents one specific sequence out of 128 different sequences that anneal to the corresponding BOX 6

neafing process.

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complementing elements.

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Reaction of S and T, followed by cleavage of the linker L leads to translocation of R2 ments of this class may anneal to both BOX 2 and 3). Reactive groups type II X and and R3 onto R4. Reaction of A with B, and C with D translocates R5 and R6 to R4. results in cleavage of R1 from the complementing element and translocation to R4. pair is orthogonal to S/T orthogonal to A/B orthogonal to C/D. Reaction of X with Y In this example, the functional entity of the monomer binding to BOX 4 serves as a Y react; S and T react; A and B react; and C and D react. In the example the X/Y (BOX 2 and 3 are of the same class, i.e., the corresponding complementing ele-(C) Templating using six monomers. Five classes of coding elements are used

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scaffold" onto which is added various substituents.

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Figure 39: A typically panning protocol for selection of templated molecules

encoding technology. These templated molecules are incubated with the immobi-Templates presenting the various small molecule variants are produced by DNA ized target molecule. Templated molecules with low affinity for the target are

amplified using PCR. The enriched templates are then ready to be used as a coding washed away. The remaining templated molecules are eluted and the template is strand for the next cycle. S

Figure 40: Array of templated molecules

- nat on a suitable surface. The templated molecule library (single-stranded template DNA) is added and allows hybridizing to the complement DNA strand. This will allow site-specific immobilization of the templated molecules. A biological sample containng target molecules is added and non-bound material is washed off. The final step The figure shows a templated molecule chip. A DNA library Is spotted in array foris the detection of bound material in each single spot. 2 5
- Figure 41. Use of rigid or partially rigid linkers to increase probability of reaction between the functional entitles of the incorporated monomer building blocks.
- (A) By using linkers comprising one or more flexible regions ("hinges") and one or more rigid regions, the probability of two functional entities getting into reactive contact may be increased. ឧ
- Symbol used for monomer building block with a rigid part and two flexible æ
- complementary sequence to the central part of the linker leads to formation of a rigid double helix; at either end of the linker a single-stranded region (C) A monomer building block with the characteristics described in (B): The element (horizontal line), and a ollgonucleotide as linker connecting the monomer building block contains an oligonucleotide as complementing functional entity (FE) with the complementing element. Annealing of a 22 ္က

remains, which constitutes the two flexible hinges.

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Figure 42. Use of zipper box to increase probability of reaction between the

functional entities of the incorporated monomer building blocks.

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(A) The linkers in this example carry zipper boxes (a) or (a'), that are complementary. By operating at a temperature that allows transient interaction of (a) and (a'), the reactive groups X and Y are brought into close proximity during multiple annealing events, which has the effect of keeping X and Y in close proximity in a larger fraction of the time than otherwise achievable. Alternatively, one may cycle the temperature between a low temperature (where the zipper boxes painwise interacts stably), and a higher temperature (where the zipper boxes are apart, but where the complementing element remains stably attached to the coding element of the template). By cycling between the high and low temperature several times, a given reactive group X is exposed to several reactive groups Y, and eventually will react to form

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(B) Sequences of two oligonucleotide-based monomer building blocks. The region constituting the complementing element, linker and zipper box is indicated.

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Figure 43. Templated synthesis of organic compounds - examples.

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(A) Three monomer building blocks are used. Each monomer building block comprises an activated ester (reactive group of type II, (X)) where the ester molety carries a functional group Rx. Upon reaction between the esters and the amines on the scaffold (scaffold may be attached to the template), amide bonds are formed, and the Rx groups are now coupled to the scaffold via amide bonds. This is thus an example of simultaneous reaction (amide formation) and activation (release of the Rx moiety from the complementing, elements), see figure 29.

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(B) Analogously to (A), three amines react with three esters to form three amide bonds, thereby coupling the functional groups Rx to the scaffold moiety. However, as opposed to (A), the scaffold is here encoded by the template.

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(C) Three monomer building blocks are used. The nucleophilic amine at the far right (part of the anchorage point) attacks the ester carbonyl of the third monomer; the amine of the third monomer attacks the thioester of the second monomer, and the Horner-Wittig Emmans reagent of the first monomer reacts with the aldehyde of the third monomer under alkaline conditions.

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This forms the templated molecule. The double bond may be posttemplating modified by hydrogenation to form a saturated bond, or altematively, subjected to a Michael addition.

(D) The thiol of the scaffold reacts with the pyridine-disulfide of monomer 1.

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The amine of the scaffold reacts with the ester of the second monomer.

The double nitril activated alpha-position is acylated by the monomer 3's thioester in the presence of base. The aryllocide undergoes Suzuki coupling with the arylboronate of monomer 4 to yield the biaryl molety.

(E) Monomer 1 acylates the primary amine. The aryliodide undergoes a Suzukl coupling by monomer 2 and the benzylic amine is acylated by monomer 3.

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Acylation of the hydrazine followed by cyclization leads to formation of an hydroxypyrazole. The arylbromide undergoes Suzuki coupling with the aryl boronate of monomer 1 and finally the aldehyde reactions with the Homer-Wittig-Emmons reagent of monomer 4 to yield an alpha, beta-unsaturated amide, which may be further functionalized by either reduction with HyPd-C or undergo Micael addition with nucleophiles.

Figure 44. α- and p-peptides, hydrazino peptides and peptoids. Encoding by use of oligonucleotide-based monomer building blocks.

20 It is shown how templated synthesis may be used to generate α- and β-peptides, hydrazino peptides and peptolds.

Figure 45. Templating of تر ، ۴ ، ۴ ، ۳ م رت و و Figure 45. Templating of تر ، ۴ ، ۴ ، ۳ ما رت ال Figure 45. drides

25 It is shown how templated synthesis may be used to generate α-, β-, γ- and α-peptides, through the use of cyclic anhydrides.

Figure 46. Generation of new reactive groups upon reaction of the reactive

30 groups X and Y.

In cases where the reaction of X and Y leads to formation of a new reactive group Z, this may be exploited to increase the diversity of the templated molecule, by incorporating monomer building blocks carrying reactive groups Q that react with Z.

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(A) X and Y react to form Z, which in itself does lead to release from the complementing element. Upon reaction of Z with Q, and cleavage of the linker that connects Z to the complementing element, the templated molecule is formed.

(B) In this case, reaction of X and Y to form Z simultaneously cleaves the linker connecting X to the complementing element. Upon reaction of Z with Q, the templated molecule is formed.

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Figure 46, example 1. Templated synthesis by generating a new reactive

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The reaction of the functional entities of the first three monomer building blocks leads to formation of two double bonds, which may react with two hydroxylamines carried in by the monomer building blocks added in the second step, and leads to formation of an ester, which may react with the an hydroxylamine, carried in by the monomer added in the second step. Finally, the linkers are cleaved, generating the templated molecule.

Figure 47. Cleavable linkers.

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Cleavable linkers, the conditions for their cleavage, and the resulting products are shown.

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Figure 48. Post-templating modification of templated molecule.

After the templating process has been performed, the templated molecules may be modified to introduce new characteristics. This list describes some of these post-templating modifications.

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Figure 49 shows the result of example 64.

Figure 50 shows the result of example 65.

Figure 51 shows the result of example 66.

Figure 52 shows the result of example 67.

Figure 53 shows the result of example 68,

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Figure 54 shows the result of example 72.

Figure 55 shows the display of a templated molecule attached to the complementing template.

Figure 56 show the result of example 99.

Figure 57 A and B show the result of example 99.

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Figure 58 A and B show the result of example 99.

Figure 59 shows the result of example 99.

Figure 60 shows the result of example 102.

Figure 62 show the result of example 105.

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Figure 61 show the result of example 104.

Figure 63 show the result of example 106.

Figure 64 show the result of example 112.

Definitions

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 α -peptide: Peptide comprising or assentially consisting of at least two α -amino adds linked to one another by a linker including a peptide bond.

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acids are those not listed in Table 2. Examples of non-natural amino aclds are those the amino group present at the amino terminal end of an amino acid or peptide, and carbon atoms, comprising at least one side chain or functional group. NH2 refers to COOH refers to the carboxy group present at the carboxy terminal end of an amino Biol. Chem., 243:3552-59 (1969) and adopted in 37 C.F.R., section 1.822(b)(2) beong to the group of amino acids listed in Table 2 herein below. Non-natural amino Amino acid: Entity comprising an amino terminal part (NH₂) and a carboxy terminal isted e.g. in 37 C.F.R. section 1.822(b)(4), all of which are incorporated herein by part (COOH) separated by a central part comprising a carbon atom, or a chaln of natural amino acids. Natural amino acids of standard nomenclature as listed in J. reference. Further examples of non-natural amino acids are listed herein below. Amino acid residues described herein can be in the "D" or or "L" isomeric form. acld or peptide. The generic term amino acid comprises both natural and non-

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ohenylatanine jutamic acid nethionine ryptophan soleucine **Jutamine** hreonine histidine alanine eucine proline serine valine ysine 1-Letter 3-Letter Met ¥ E P da 딩 Ala Ser Ľ ᢟ 윤 횬 ₽ Symbols ≥ ∞ αш I 2 23 ဓ

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aspartic acid asparagine cysteine Asn Asp Š

Table 2. Natural amino aclds and their respective codes. S

Amino acid precursor: Moiety capable of generating an amino acid residue following ncorporation of the precursor into a peptide.

- such reactions known in the art can be used as appropriate as readily recognized by hose skilled in the art. Accordingly, templated molecules can be amplified by using he polymerase chain reaction (PCR), ligase chain reaction (LCR), in vivo amplifica-Amplifying: Any process or combination of process steps that increases the number carried out by any state of the art method including, but not limited to, a polymerase sule being templated by the template. Any amplification reaction or combination of tion of cloned DNA, and the like. The amplification method should preferably result plates for synthesising additional copies of the templated molecules comprising a sequence of functional groups resulting from the synthesis of the templated moleproportions of templates of different sequences in a mixture prior to amplification. chain reaction to increase the copy number of each template, and using the temn the proportions of the amplified mixture being essentially representative of the of copies of a templated molecule. Amplification of templated molecules may be 9 5 8
- Base: Nitrogeneous base molety of a natural or non-natural nucleotide, or a derivalive of such a nucleotide comprising alternative sugar or phosphate moleties. Base capable of complementing one or more bases of the opposite nucleotide strad of a moleties Include any molety that is different from a naturally occurring molety and double helix.

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- coding element, b) at least one functional entity comprising a functional group and a reactive group, and c) at least one linker separating the at least one functional entity from the at least one complementing element, wherein the building block does not comprising at least one recognition group capable of recognising a predetermined Building block: Species comprising a) at least one complementing element ဧ
 - comprise a ribosome. Preferred building blocks are capable of being incorporated 32

arginine

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into a nucleotide strand and/or capable of being linked by reactions involving reactive groups of type I and/or type II as described herein. Cleavable linker: Residue or bond capable of being cleaved under predetermined conditions.

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Cleaving: Breaking a chemical bond. The bond may be a covalent bond or a noncovalent bond.

recognition may result from the formation of a covalent bond or from the formation of Coding element: Element of a template comprising a recognition group and capable of recognising a predetermined complementing element of a building block. The a non-covalent bond between corresponding pairs of coding elements and complementing elements capable of interacting with one another.

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predetermined complementing element capable of recognising said coding element. Coding element complementation: Contacting a coding element with a

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nucleotide comprising a base moiety, predetermined sets of nucleotides are capable predetermined complementing element capable of recognising said coding element. Complementing: Process of bringing a coding element into reactive contact with a of complementing each other by means of hydrogen bonds formed between the When the coding element and the complement element comprises a natural base moleties.

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Complementing element: Element of a building block. Linked to at least one functional entity by means of a linker. See coding element.

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Complementing template: A sequence of complementing elements, wherein each element. A complementing element is capable of recognising a predetermined complementing element is covalently linked to a neighbouring complementing coding element. The complementing template may be linear or branched.

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Complex: Templated molecule linked to the template that templated the synthesis of the templated molecule. The template can be a complementing template as defined

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herein that is optionally hybridised or otherwise attached to a corresponding

template of linked coding elements.

partners or hybridization partners into reactive contact with each other. The reactive Contacting: Bringing e.g. corresponding reactive groups or corresponding binding contact is evident from a reaction or the formation of a bond or a hybridization between the partners. S

Corresponding binding partners: Binding partners capable of reacting with each

other. 2 Corresponding reactive groups: Reactive groups capable of reacting with each other.

comprises a functional group and a reactive group capable of linking neighbouring, Functional entity: Entity forming part of a building block. The functional entity functional groups. 5

functional groups in a templated molecule is a result of the capability of the template Functional group: Group forming part of a templated molecule. The sequence of to template the synthesis of the templated molecule.

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mediated by recognition groups forming corresponding binding partners by means of complementing elements into reactive contact with each other. The reaction may be emplate comprising a plurality of coding elements with a plurality of building blocks. covalent or non-covalent bonds. The interaction may occur as a result of mixing a nteracting: Used interchangably with contacting. Bringing species such as e.g. correspnding binding partners in the form of e.g. coding elements and

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Ligand: Used herein to describe a templated molecule capable of targeting a target there can exist more than one ligand for a given target. The ligands can differ from molecule. In a population of candidate template molecules, a ligand is one which oinds with greater affinity than that of the bulk population. In a candidate mixture one another in their binding affinities for the target molecule. င္က

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allow the species some freedom of movement in relation to each other. The link can may be retained at an essentially fixed distance, or the linker may be flexible and Linker: A residue or chemical bond separating at least two species. The species complementing element and a functional entity of a building block, neighbouring complementing template, and neighbouring functional groups of a templated coding elements of a template, neighbouring complementing elements of a be a covalent bond or a non-covalent bond. Linked species include e.g. a

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phates of ribo or deoxyribo derivatives of A, T, U, C, or G) are directed by a template system composed of a fused, six membered ring and a five membered ring (a purine in natural oligonucleotides), with a middle hydrogen bond linking two ring atoms, and bonds formed between the bases. In the geometry of the Watson-Crick base pair, a enzymes are able to catalyze the synthesis of an oligonucleotide complementary to the template ollgonucleotide. In this synthesis, building blocks (normally the triphosconsists of a sugar moiety (ribose or deoxyribose), a phosphate moiety, and a natucleotide strands. The base pairing results in a specific hybridization between prede-Natural nucleotide: Any of the four deoxyribonucleotides, dA, dG, dT, and dC (con-RNA) are the natural nucleotides. Each natural nucleotide comprises or essentially (C), wherein corresponding base-pairs are part of complementary, anti-parallel nubase pairs. In nature, the specific Interactions leading to base pairing are governed by the size of the bases and the pattern of hydrogen bond donors and acceptors of the bases. A large purine base (A or G) pairs with a small pyrimidine base (T, U or six membered ring (a pyrimidine in natural oligonucleotides) is juxtaposed to a ring (A) pairs with thymine (T) or uracil (U); and where guanine (G) pairs with cytosine according to well-known rules of base pairing (Watson and Crick), where adenine termined and complementary nucleotides. The base pairing is the basis by which oligonucleotide to form a complementary oligonucleotide with the correct, complementary sequence. The recognition of an oligonucleotide sequence by its compleral/standard base moiety. Natural nucleotides bind to complementary nucleotides C). Additionally, base pair recognition between bases is influenced by hydrogen hydrogen bonds on either side joining functional groups appended to each of the stituents of DNA), and the four ribonucleotides, A, G, U, and C (constituents of mentary sequence is mediated by corresponding and interacting bases forming rings, with donor groups paired with acceptor groups. 2 5

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mentioned complementing elements are sald to be neighbouring and sald two com-Nelghbouring: Elements, groups, entities or residues located next to one another In plementing elements define neighbouring functional entitles and neighbouring codments, each linked to a functional entity, are linked to one another through one (or more) complementing element(s) that is not linked to a functional entity, the aforea sequence are said to be neighbouring. In cases where two complementing eleing elements that can be linked to one another, either directly or through one (or

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more) coding element(s).

Non-natural amino acid: Any amino acid not included in Table 2 herein above. Nonnatural amino acids includes, but is not limited to modified amino acids, L-amino acids, and stereoisomers of D-amino acids. Non-natural base pairing: Base pairing among non-natural nucleotides, or among a 5,037,120, wherein eight non-standard nucleotides are described, and wherein the through which the base pairing Is established are different from those found In the natural AT, AU and GC base pairs. In this expanded set of base pairs obeying the Matson-Crick hydrogen-bonding rules, A pairs with T (or U), G pairs with C, iso-C natural base has been replaced by a non-natural base. As is the case for natural pairing with a fused, bicyclic heterocyclic ring system composed of a five member pairs with iso-G, and K pairs with X, H pairs with J, and M pairs with N (Figure 2). nucleotides, the non-natural base pairs involve a monocyclic, six membered ring natural nucleotide and a non-natural nucleotide. Examples are described in US Ing fused with a six membered ring. However, the patterns of hydrogen bonds 15 ನ 22

bonding rules have also been described (Berger et al., 2000, Nucleic Acids Re-Nucleobases capable of base pairing without obeying Watson-Crick hydrogensearch, 28, pp. 2911-2914). Non-natural nucleotide: Any nucleotide not falling within the definition of a natural nucleotide. ႙

Into an oligonucleotide, preferably by means of an enzyme comprising DNA or RNA natural nucleotides capable of being incorporated - in a template-directed manner -Nucleotide: Nucleotides as used herein refers to both natural nucleotides and non-

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comprising a nucleotide part are capable of Interacting with each other by means of may differ from natural nucleotides by having a different phosphate moiety, sugar equivalents of natural or recombinant DNA or RNA polymerases. Corresponding hydrogen bonds. The Interaction is generally termed "base-pairing". Nucleotides bond in the form of a phosphodiester bond, or in the form of a non-natural bond, respective neighbour(s) in a template or a complementing template by a natural dependent DNA or RNA polymerase activity, including variants and functional binding partners in the form of coding elements and complementing elements molety and/or base molety. Nucleotides may accordingly be bound to their such as e.g. a peptide bond as in the case of PNA (peptide nucleic acids).

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emplate. Nucleotide analogs often Includes monomers or oligomers containing noncleotide analogs includes e.g. nucleotides in which the phosphodiester-sugar backother monomers and/or oligomers through specific base pairing is possible. Alternastrate of enzymes, such as DNA polymerases and RNA polymerases, or mutants or natural bases or non-natural backbone structures that do not facilitate incorporation Nucleotide analog: Nucleotide capable of base-pairing with another nucleotide, but incapable of being incorporated enzymatically into a template or a complementary functional equivalents thereof, are defined as nucleotide analogs herein. Oligonubone of natural oligonucleotides has been replaced with an alternative backbone tive oligomers capable of specifically base paining, but unable to serve as a subinto an oligonucleotide in a template-directed manner. However, interaction with nclude peptide nucleic acid (PNA), locked nucleic acid (LNA), and morpholinos.

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appended molecular entity. Often, derivatized building blocks (nucleotides to which oligonucleotides by RNA or DNA polymerases, using as substrate the triphosphate nucleotides can be derivatized on the bases, the ribose/deoxyribose unit, or on the phosphate. Preferred sites of derivatization on the bases include the 8-position of adenine, the 5-position of uracil, the 5- or 6-position of cytosine, and the 7-position that the derivative is inserted opposite a predetermined nucleotide in the template. incorporated Into the growing oligonucleotide chain with high specificity, meaning a molecular entity have been appended) can be enzymatically incorporated into of the derivatized nucleoside. In many cases such derivatized nucleotides are Nucleotide derivative: Nucleotide or nucleotide analog further comprising an Such an incorporation will be understood to be a specific incorporation. The

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corresponding positions (Benner, United States Patent 6,037,120). Other sites of of guanine. The nucleotide-analogs described below may be derivatized at the pairing specificity. Preferred sites of derivatization on the ribose or deoxyribose derivatization may be used, as long as the derivatization does not disrupt base

- stabilize the nucleic acids towards degradation, and it may be advantageous to use phosphates may be derivatized. Preferred derivatizations are phosphorothiote. moleties are the 5', 4' or 2' positions. In certain cases it may be desirable to employed, as long as the base pairing specificity is not disrupted. Finally, the 2'-modified nucleotides (US patent 5,958,691). Agaln, other sites may be Ŋ
- It is clear that the various types of modifications mentioned herein above, including i) Nucleotide analogs (as described below) may be derivatized similarly to nucleotides. derivatization and ii) substitution of the natural bases or natural backbone structures respectively, can be applied once or more than once within the same molecule. with non-natural bases and alternative, non-natural backbone structures, 2

nucleotides, including any combination thereof. The natural and/or non-natural nucleotides may be linked by natural phosphodiester bonds or by non-natural oligonucleotide comprises oligonucleotides of both natural and/or non-natural Oligonucleotide: Used herein Interchangebly with polynucleotide. The term bonds. Oligonucleotide is used interchancably with polynucleotide.

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Oligomer: Molecule comprising a plurality of monomers that may be Identical, of the same type, or different. Oligomer is used synonymously with polymer in order to describe any molecule comprising more than two monomers. Oligomers may be

- different monomers of the same type, or heterooligomers comprising different types homooligomers comprising a plurality of identical monomers, oligomers comprising of monomers, wherein each type of monomer may be identical or different. 25
- linked to a template, that do not have an affinity for and is consequently not bound Partitioning: Process whereby templated molecules, or complexes comprising such to - such target molecules. Partitioning can be accomplished by various methods molecules linked to a template, are preferentially bound to a target molecule and separated from templated molecules, or complexes comprising such molecules known in the art. The only requirement is a means for separating targeted, ဓ
 - templated molecules bound to a target molecule from templated molecules not 32

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bound to target molecules. The choice of partitioning method will depend on properties of the target molecule and of the templated molecule and can be made according to principles and properties known to those of ordinary skill in the art.

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Peptide: Plurality of covalently linked amino acid residues defining a sequence and linked by amide bonds. The term is used analogously with oligopeptide and polypeptide. The amino acids may be both natural amino acids and non-natural amino acids, including any combination thereof. The natural and/or non-natural amino acids may be linked by peptide bonds or by non-peptide bonds. The term peptide also embraces post-translational modifications introduced by chemical or enzyme-catalyzed reactions, as are known in the art. Such post-translational modifications can be introduced prior to partitioning, if desired. Amino acids as specified herein will preferentially be in the L-stereolsomeric form. Amino acid analogs can be employed instead of the 20 naturally-occurring amino acids. Several such analogs are known, including fluorophenylalanine, norleucine, azeitdine-2-carboxylic acid, S-aminoethyl cysteine, 4-methyl tryptophan and the like.

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Plurality: At least two.

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Polymer: Templated molecule characterised by a sequence of covalently linked residues each comprising a functional group, including H. Polymers according to the invention comprise at least two residues.

Polynucleotide: See oligonucleotide,

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Precursor: Molety comprising a residue and being capable of undergoing a reaction during template directed synthesis of a templated molecule, wherein the residue part of the precursor is built into the templated molecule.

Reactive group: Corresponding reactive groups being brought into reactive contact with each other are capable of forming a chemical bond linking e.g. a coding element and its complementing element, or coupling functional groups of a templated molecule.

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Recognition group: Part of a coding element and involved in the recognition of the complementing element capable of recognising the coding element. Preferred recognition groups are natural and non-natural nitrogeneous bases of a natural or non-natural nucleotide.

Recombine: A recombination process recombines two or more sequences by a process, the product of which is a sequence comprising sequences from each of the two or more sequences. When involving nucleotides, the recombination involves an exchange of nucleotide sequences between two or more nucleotide molecules at sites of identical nucleotide sequences, or at sites of nucleotide sequences that are not identical, in which case the recombination can occur randomly. One type of recombination among nucleotide sequences is referred to in the art as gene

Repetitive sequence: Sequence of at least two elements, groups, or residues, occurring more than once in a molecule.

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Residue: A polymer comprises a sequence of covalently linked residues, wherein each residue comprises a functional group.

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Ribose derivative: Ribose moiety forming part of a nucleoside capable of being enzymatically incorporated into a template or complementing template. Examples include e.g. derivatives distinguishing the ribose derivative from the riboses of natural
ribonucleosides, including adenosine (A), guanosine (G), uridine (U) and cytidine
(C). Further examples of ribose derivatives are described in e.g. US 5,788,481. The

(C). Further examples of ribose derivatives are described in e.g. US 5,786,461. The term covers derivatives of deoxyriboses, and analogously with the above-mentioned disclosure, derivatives in this case distinguishes the deoxyribose derivative from the deoxyriboses of natural deoxyribonucleosides, including deoxyadenosine (dA), deoxydyuanosine (dG), deoxydhymidine (dT) and deoxycytidine (dC).

Selectively cleavable linker: Selectively cleavable linkers are not cleavable under conditions wherein a cleavable linker is cleaved. Accordingly, it is possible to cleave the cleavable linkers linking complementing elements and functional groups in a templated molecule without at the same time cleaving selectively cleavable linkers linking - in the same templated molecule - a subset of complementing elements and

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templated molecule, wherein the template and the templated molecule are linked by molecule and the template that has directed the template-mediated synthesis of the functional groups. It is thus possible to obtain a complex comprising a templated one or more, preferably one, selectively cleavable linker(s).

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selection conditions that demand a requisite specificity, or by tailoring and modifying corresponding binding partners. Specific recognition is a term which is defined on a measured between the target molecule and a candidate template molecule mixture templated molecule as a whole and target as a whole. The templated molecules of should be comparable to the conditions of the Intended use. For the most accurate affinity of a coding element recognition group for a complementing group results in interaction of templated molecule and target molecule of a higher affinity than that reactions must be essentially similar and preferably the same, and the conditions the invention can be selected to be as specific as required, either by establishing is observed. In order to compare binding affinities, the conditions of both binding Specific recognition: The interaction of e.g. a coding element with preferably one case-by-case basis. In the context of a given interaction between predetermined the formation of predominantly only one type of corresponding binding partners. predetermined complementing element. A specific recognition occurs when the comparisons, measurements will be made that reflect the interaction between binding partners, e.g. a templated molecule and a target molecule, a binding Simple mis-match incorporation does not exclude a specific recognition of the templated molecules.

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Subunit: Monomer of coding element comprising at least one such subunit. 22

Support: Solid or semi-solid member to which e.g. coding elements can be attached Functional molecules or target molecules may also be attached to a solid support during targeting. Examples of supports includes planar surfaces including sillcon during interaction with at least one complementing element of a building block. wafers as well as beads.

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Tag: Entity capable of identifying a compound to which it is associated.

Target molecule: Any compound of interest for which a templated molecule in the 35

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transition state analog, cofactor, inhibitor, drug, controlled substance, dye, nutrient, polysaccharide, glycoprotein, hormone, receptor, receptor ligand, cell membrane form of a ligand is desired. A target molecule can be a protein, fusion protein, component, antigen, antibody, virus, virus component, substrate, metabolite, peptide, enzyme, nucleic acid, nucleic acid binding protein, carbohydrate,

(complementing) temptate of complementing elements unless otherwise specified. emplate: Template refers to both a template of coding elements and a

growth factor, toxin, lipid, glycolipid, etc., without limitation.

the templated molecule, and the templating activity involves the formation of specific When referring to a template of coding elements, each coding element is covalently wherein the complementing element forms part of a building block also comprising or branched. A template of coding elements actively takes part in the synthesis of recognising a predetermined complementing element. The template may be linear comprises a string of nucleotides, the nucleotides may be natural or non-natural, pairing partners in the form of coding element complementing element hybrids, the functional group forming part of the templated molecule. The template is inked to a neighbouring coding element. Each coding element is capable of preferably a string of nucleotides or nucleotide analogs. When the template **e** 5

analogs. The sugar moiety of a nucleotide or nucleotide analog may be a ribose or a equivalent means capable of linking nucleotide analogs so as to allow the nucleotide and may be linked by e.g. phosphorothloate bonds or natural phosphodiester bonds. analog string to hybridize specifically with another string of nucleotides or nucleotide template or complementing template to hybridise specifically to another string of Nucleotide analogs may be linked e.g. by amide bonds, peptide bonds, or any deoxyribose, a ribose derivative, or any other molecular molety that allows the nucleotides or nucleotide analogs.

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wherein the formation of a templated molecule comprising a sequence of covalently coding element of the template directs the incorporation of the functional group into incorporation and templated synthesis. Template directed synthesis is the process, relationship between coding elements and functional groups, and the contacted linked, functional groups involves contacting a string of coding elements with particular complementing elements. The process thus defines a one-to-one Template directed synthesis: Used synonymously with template directed ജ 33

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between the bases, where the bases may be natural (i.e. A, T, G, C, U), and/or noncomprises or essentially consists of nucleotides, a template directed synthesis of an in the template in a one-base-to-one-base pairing manner. The interaction specifies the incorporation of complementing nucleotides opposite their base pairing partners oligonucleotide is based on an interaction of each nucleotide with its paining partner oligonucleotide strand interact when forming specific base-pairs. This base pairing groups. Accordingly, there is a predetermined one to one relationship between the in the template. Consequently, one base, including a heterocyclic base, from each herein by reference. Further examples of non-natural bases are e.g. PNA (peptide means than hydrogen bonding (e.g. Interaction between hydrophobic nucleobases with "complementary" structures; Berger et al., 2000, Nucleic Acids Research, 28, specificity may be achieved through Watson-Crick hydrogen-bonding interactions complementing element, or otherwise - the coding element capable of temptating another nucleotide or a predetermined subset of nucleotides, for example A base pp. 2911-2914). The Interacting oligonucleotide strands as well as the individual that particular functional group into the templated molecule. When the template the templated molecule comprising a sequence of covalently linked, functional sequence of functional groups of the templated molecule and the sequence of natural bases such as those e.g. disclosed e.g. in US 6,037,120, incorporated oligonucleotides containing non-standard base pairs can be achieved by other coding elements of the template that templated the synthesis of the templated between oligomers results from the specific base pairing of a nucleotide with molecule. Thus, during the templated synthesis of the templated molecule, a functional group is initially contacting - by means of a linker moiety and/or a nucleotides are said to be complementary. The specificity of the interaction nucleic acid), LNA (lock nucleic acid) and morpholinos. Base pairing of pairing with U, and C base pairing with G. S 2 5 ೪ 32

Templated: Feature of the templated molecule of the complex comprising a template linked to the templated molecule, wherein the templated molecule is obtainable by template directed synthesis using the template. Thus, one component of the complex (the template) is capable of templating the synthesis of the other component (the templated molecule). The term is also used to describe the synthesis of the templated molecule that involves the incorporation into the templated molecule of functional groups, wherein the incorporation of each

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functional group involves contacting a coding element with a particular functional group, or with a building block comprising said functional group, wherein the contacted coding element of the template directs the incorporation of functional groups into the templated molecule linked to the template that templated in this way the synthesis of the templated molecule. Thus, during the templated synthesis of the templated molecule, a functional group is initially contacting - either directly or by means of a linker molety and/or a complementing element - the coding element capable of templating that particular functional group into the templated molecule.

10 Templated molecule: Molecule comprising a sequence of covalently finked.

functional groups, wherein the templated molecule is obtainable by template directed synthesis using the template. Thus, one component of the complex (the template) is capable of templating the synthesis of the other component (the templated molecule). When the template comprises or essentially consists of nucleotides, the template is capable of being amplified, wherein said template amplification results in a plurality of templated molecules, wherein each templated molecule is generated by template directed synthesis using the templated molecules can be generated by a template directed synthesis using either a template of coding elements or a complementing template, template of coding elements or a complementing template of complementing elements as a template for the template directed synthesis of the templated molecule.

Templating: Process of generating a templated molecule.

25 Variant: Template or templated molecule exhibiting a certain degree of identity or homology to a predetermined template or templated molecule, respectively.

Detailed Description of the Invention

In one preferred embodiment of the present invention, there is provided a "chemical display of templated molecules" which enables the generation of a huge number of "templated polymers" (e.g. from about 10³ to about or more than e.g. 10¹⁸ as described elsewhere herein), wherein each templated molecule is individually linked to a "template" that serves as identification of that individual polymer (its sequence of

residues), as well as a means for amplification (many copies of the molecule can be

prepared by a process that replicates the template). Preferred embodiments of the invention are disclosed in Figure 1 illustrating various steps of the method of the Invention.

Step 1. Synthesis

which have been appended a functional entity through a cleavable linker, and where functional entity and a complementing element that are linked by means of a cleavthe functional entity comprises or essentially consists of an "activatable" polymer able linker (Figures 3 and 4). Preferred building blocks comprise a nucleotide to Different monomer building blocks are synthesized. Building blocks comprise a unit (Figure 6).

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Step 2. Incorporation

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formed, from which the functional entitles protrude. The sequence of functional entigonucleotide strand according to the directions of a oligonucleotide template. As a iles is determined by the sequence of coding elements, such as nucleotides, of the polymerase is preferably used to incorporate the nucleotide-derivatives into an oli-The building blocks are used as substrates in a template-dependent polymer synthesis. In one embodiment, the building blocks are nucleotide-derivatives and a result, a complementing template (a string of incorporated building blocks) is template.

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encoding portion of the template (see for example Figure 7 or 8), or the link could be linker which, after polymerization and activation, is capable of linking the templated Figure 1 describes the use of a building block that carries the selectively cleavable polymer to its template. Alternatively, the selectively cleavable linker can be comprised by an oligo capable of annealing upstream or downstream of the polymerto the template directly.

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The building block can preferably be incorporated by an enzyme, such as for examsuch as mono-, di- or polynucleotides. In some of these cases, a primer is required ple DNA polymerase, RNA polymerase, Reverse Transcriptase, DNA ligase, RNA ligase, Taq DNA polymerase, HIV-1 Reverse Transcriptase, Klenow fragment, or any other enzyme that will catalyze the incorporation of complementing elements

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(for example DNA polymerase). In other cases, no primer is required (e.g., RNA polymerase).

Step 3. Polymerization

Each functional entity has preferably reacted with neighbouring functional entities to form a polymer during or after formation of the complementing template. A change In conditions, e.g., photolysis, change in temperature, or change in pH, may initiate the polymerisation either during or after complementing template formation. ιO

Step 4. Activation 2

cleavable under conditions resulting in cleavage of the remaining linkers. The result more predetermined position(s), including a single position, where the linker is not cleavage of at least one linker, or a plurality of cleavable linkers, except at one or The formed polymer is preferably released from the complementing elements by Is a templated polymer attached at one or more positions, preferably only at one

Step 5. Selection and amplification

position, to the template that encodes it.

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characteristics (e.g., binding to a receptor) are recovered and amplified, by first am-A selection process can subsequently be performed, wherein a huge number of difsis, is challenged with a molecular or physical target (e.g. a biological receptor or a ferent templated molecules, each attached to the template that directed its synthesurface), or is exposed to a certain screen. Templated molecules having desired plifying the templates, and then using the templates for a new round of templated 8

polymer synthesis. The process of selection and amplification can be repeated several times, until a polymer with appropriate characteristics (e.g., high affinity for the receptor) is isolated. 23

templated molecule complexes with affinity for the immobilized target molecule. The molecular target (e.g., a receptor) had been immobilized. After washing the column, the binders are eluted. This eluate consists of an enriched population of templatetemplate-templated molecule complexes to an affinity column, to which a certain subjected to yet a selection round, where the conditions optionally may be more enriched population may be taken through an amplification round, and then be A typical selection protocol involves the addition of a population (a library) of ജ

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stringent. After a number of such selection-and-amplification rounds, an enriched population of high affine binders are obtained.

When selecting for the ability of a templated molecule to become internalized into a cell, the selection step may involve a simple mixing of the population of template-templated molecule complexes with cells. After incubation (to allow the internalization of the template-templated molecule complexes), the cells are washed, and the internalized template-templated molecule complexes may be recovered by lysis of the cells. As above, the template-templated molecule complexes may be amplified and taken through further rounds of selection-and-amplification. After a number of selection-and-amplification rounds, an enriched population of templated molecules with the ability to internalize are obtained.

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Building blocks - molecular design

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The building blocks (also termed "monomers") is preferably of the general design shown in Figure 3 and 4. The monomer in one embodiment coriprises the following elements: Complementing element-Linker-Backbone comprising reactive group(s) type II-Functional group, where the complementing element comprises or essentially consists of a recognition group and reactive group(s) type I. In this case the linker is preferably a "traceless linker", i.e., a linker that does not leave any (undesirable) molecular entity on the functional entity. Building blocks with this composition are used in for example (Figure 15, example 7).

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Alternatively, the monomer may have the composition Complementing elsment-Linker-Functional Group-Backbone containing reactive group(s) type II, in which case the desired functional group is created as a result of cleavage of the linker. Building blocks with this composition are used in for example (Figure 17, example 1).

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The functional groups must be compatible with the desired method for incorporation of complementing elements, their polymerization and activation. Obviously, it is important to preserve the integrity of the template and the templated molecule in these processes.

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Functional groups that are not compatible with the conditions of incorporation, polymerization or activation must be protected during these processes, or alternatively, the functional groups must be introduced after these processes have taken place.

The latter is done by templating a functional group (e.g., an activated disulfide) that is compatible with the incorporation, polymerization and activation, and that will specifically react with a bifunctional molecule (e.g., a thiol connected to the desirable functional group, R_k), added after activation. Alternatively, functionalities may be introduced by e.g. oxidation, or any other form of treatment, of the incorporated functional entities after activation. In this way, functionalities such as components of natural effector molecules or synthetic drugs that are otherwise difficult to handle,

In some embodiments of the process of the invention as described herein, there is no need for a cleavable linker, as the polymerisation reaction involves cleavage of the linker (Figure 14 and Figure 14, example 1).

may be incorporated.

When being nucleotides, the complementing elements may contain one, two or several nucleotides or nucleotide-analogs. The use of di-, tri- or longer oligonucleotides presents a number of advantages. First, a higher monomer diversity may be encoded by the template. Second, the requirements for the site of attachment of the functional entity to the complementing element becomes more relaxed. Third, there would be less bulk per mononucleotide in the formed polynucleotide, potentially leading to higher display-efficiencies. Fourth, it would allow the display of polymers with longer residue-unit-length. Also, it would allow the display of bigger functional

In cases where a polymerase is employed for the incorporation of nucleotide comprising building blocks, it is preferred that the nucleotides are derivatized in a way that allows their specific and efficient incorporation into the growing strand.

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More than 100 different nucleoside- and nucleotide-derivatives are commercially available or can be made using simple techniques (Eaton, Current Opinion in Chemical Biology, 1897, 1: 10-16). Moreover, many nucleotide-derivatives, modified on the bases or the riboses, are incorporated efficiently and specifically by varified on the bases or the riboses, are incorporated efficiently and specifically by varified on the bases or the riboses, are incorporated efficiently and specifically by varified on the bases or the riboses.

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(Figure 9). Nucleotides with additions of up to 300 Da have been incorporated specifically and efficiently (Wiegand et al., Chemistry and Biology, 1997, 4: 675-683; Fenn and Herman, Analytical Chemistry, 1990, 190: 78-83; Tarasow and Eaton, ous polymerases, in particular T7 RNA polymerase and Reverse Transcriptase

Biopolymers, 1998, 48: 29-37). In addition to the four natural base pairs (AT or AU, TA or UA, CG, GC), at least 8 base pairs are known to hybridise specifically, some of which are incorporated into oligonucleotides by polymerases in a templatedependent manner.

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T7 RNA polymerase and T7 RNA polymerase mutant Y639F, Sequenase, Taq DNA T7 DNA polymerase, T4 DNA polymerase, T4 DNA Ligase, E. coli RNA polymerase, cleotides and/or oligonucleotides as substrates. Mutant or engineered polymerases The incorporation of complementing elements may be catalyzed by chemical or biopolymerase, Klenow Fragment (Large fragment of DNA polymerase I), DNA-ligase, merase, ribozymes with ligase or replicase activities such as described in (Johnston transcriptases, and DNA- and RNA- ligases, ribozymes and deoxyribozymes. Spewith improved characteristics, for example broadened nucleotide substrate specificcific examples include HIV-1 Reverse Transcriptase, AMV Reverse Transcriptase, rTh DNA polymerase, Vent DNA polymerase, Pfu DNA polymerase, Tte DNA polyet al., Science, May 18, 2001, pp. 1319-1325), and other enzymes that accept nucatalysts are template-dependent DNA- and RNA-polymerases, including reverse logical catalysts. When the building blocks are nucleotides, particularly relevant 2 5

use single or double stranded nucleotides as templates, and produce single or douby deleting the nuclease activity), are particularly relevant. The polymerases may ble stranded nucleotide products.

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Sites of modification that have been shown to be accepted by polymerases include

the following non-exhaustive list of examples (See also Figure 9):

ity, and mutants in which the proofreading function has been eliminated (for example

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Site of modification 3-position 7-position Nucleotide dATP dATP

8-position dATP dATP

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2' (deoxyribose molety)

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4' (deoxyribose moiety) dTP

7-position dGTP

2' (deoxyribose moiety) dCTP

2' (deoxyribose) dCTP

5-position 5

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8-position

ATP

Terminal transferase, RNA ligases, Polynucleotide kinases and other template indeincluding engineered or mutant variants, may be used for some of the applications pendent enzymes that accept nucleotides and/or oligonucleotides as substrates,

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and method variations described in the present invention.

ploying complementing elements that are di-, tri- or polynucleotides, it may be possible to attach functional entitles at these alternative sites without inhibiting specific without eliminating hybridization or Incorporation specificity. Particularly when em-It may be possible to attach the functional entitles at other sites in the nucleotide, ncorporation. ន

Cleavable and non-cleavable linkers

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cleave them, are illustrated in (Figure 10). In one aspect of the invention, the linker A selection of cleavable linkers and protection groups, as well as the agents that may be selected from the following list: Carbohydrides and substituted carbo-

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Aryl/hetaryl, polyaryl/hetaryl and substituted polyaryl/polyhetaryl; Ethers, polyethers such as e.g. polyethylenglycol and substituted polyethers; Amines, polyamines and hydrides; Vinyl, polyvinyl and substituted polyvinyl; Acetylene, polyacetylene; substituted polyamines; Double stranded, single stranded or partially double

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stranded natural and unnatural polynucleotides and substituted double stranded, single stranded or partially double stranded natural and unnatural polynucleotides; Polyamides and natural and unnatural polypeptides and substituted polyamides and natural and unnatural polypeptides.

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It one aspect of the invention it is preferred that linkers do not react with other linkers, complementing elements or functional entities, in the same monomer or in another monomer. Also, in some of the schemes proposed herein, it is desirable that the linker is not cleaved by the conditions of polymerization. Finally, it is preferred that the conditions of linker cleavege does not affect the integrity of the template, complementing template or functional entitles.

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Linkers can be cleaved in any number of ways when subjected to predetermined conditions. Linkers may e.g. be cleaved with acid, base, photolysis, increased temperature, added agents, enzymes, ribozymes or other catalysts. Examples of cleavable linkers and their respective protection groups are shown in (Figure 10), along with the conditions for linker cleavage, and the cleavage products.

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To maintain a physical link between the template and the templated molecule, at least one non-cleavable linker is needed. This non-cleavable linker is preferably flexible, enabling it to expose the templated molecule in an optimal way.

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Functional groups

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The one or more functional groups that appear on the functional entity may be selected from a variety of chemical groups which gives the templated molecules the desired properties or serves another beneficial purpose, like higher lipophilicity for recovery purposes. A non-limiting selection of functional groups is indicated below: Hydroxy; alkoxy; Hydrogen; Primary, secondary, tertiary amines; Carboxylic acids esters; Phosphates, phosphonates; Sulfonates, sulfonamides; Amides; Carbomates; Carbomates; Ureas; Alkenes, Alkenes, Alkynes; Anhydrides; Katones; Aldehydes; Nitatrates, nitrites; Imines; Phenyl and other aromatic groups; Pyridines, pyrimidines, purines, indole, Imidazole, and heterocyclic bases; Heterocy-

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Small molecule catalysts; Dextrins, saccharides; Fluorescein, Rhodamine and other

cles; polycycles; Flavins; Halides; Metals; Chelates; Mechanism based inhibitors;

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fluorophores; Polyketides, peptides, various polymers; Enzymes and ribozymes and other biological catalysts; Functional groups for post-polymerization/post activation coupling of functional groups; Drugs, e.g., taxol moiety, acyclovir moiety, "natural products"; Supramolecular structures, e.g. nanoclusters; Lipids; and Oligonucleo-

tides, oligonucleotide analogs (e.g., PNA, LNA, morpholinos).

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Reactive groups of type II

A variety of reactive groups II may be used in the templated synthesis. Examples of reactive groups include, but are not limited to N-carboxyanhydrides (NCA), N-

10 thiocarboxyanhydrides (NTA), Amines, Carboxylic acids, Ketones, Aldehydes, Hydroxyls, Thiols, Esters, Thioesters, conjugated system of double bonds, Alkyl halides, Hydrazines, N-hydroxysuccinimide esters, Epoxides, Haloacetyls, UDP-activated saccharides, Sulfides, Cyanates, Carbonylimidazole, Thiazinanones, Phosphines, Hydroxylamines, Sulfonates, Activated nucleotides, Vinylchloride, Alkenes, and quinines.

Polymerization

Reactions that lead to polymer formation are termed polymerization reactions. The major reaction-classes are anionic polymerizations, cationic polymerizations, and pericyclic polymerizations.

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Although polymerisation reactions in solution is achievable by state of the art methods, polymerisation of functional entities linked to an array as described herein does not constitute standard type reactions. Only a few polymerisation reactions have so far been performed in an array format, and not in connection with the methods of the present invention. Consequently, it will be a matter of molecular design of the func-

- far been performed in an array format, and not in connection with the methods of the present invention. Consequently, it will be a matter of molecular design of the functional entities and their linkers and attachment points on the complementing elements (e.g. attachment to the base, ribose or phosphate of a nucleotide), as well as a matter of optimising the polymerisation conditions, in order to preferably reduce minimize or even eliminate any undesirable reactions taking place in solution while increasing or maximizing a correct template-directed polymerisation on the array.
- The present invention in one embodiment employs polymertzation reactions which are in principle known from the state of the art in the sense that they are routinely

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used in solution synthesis schemes. However, in the present invention, the reactents (reactive groups) are held in close proximity by their attachment to elements of a complementing template. This increases the local concentration significantly. Typical synthesis schemes in solution use 1 µM ~ 1 mM concentrations of the reac-

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tants. When arrayed as disclosed herein, the local concentration will typically be from a thousand-fold to a million-fold higher. As a result, the reactions can in principle be much more efficient. However, the reactions are preferably designed in such a way that the occurrence of undesirable slde-reactions are avoided. The molecular design and the polymerization conditions according to the invention reflect this fact and can be further optimised by the skilled person searching for the polymerization conditions and molecular design that maximizes the relative template directed polymerization in solution.

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Depending on the type of initiator and reactive groups, the polymerization may be initiated and/or catalyzed by changes in pH and/or temperature, addition of reactants or catalysts, enzymes or ribozymes, or light, UV or other electromagnetic radiation, etc. Particularly relevant enzymes include proteases, protein figase (e.g., subtiligase), UDP-glycogen synthetases, CGTases and polyketide synthases. In cases where the conditions and molecular designs have been finely adjusted, so as to allow efficient polymerization of the reactants when arrayed on the complementing template, but insignificant reaction in solution, the polymerization need not be initiated. The increased local concentration in the array simply drives the polymerization.

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In the case where incorporation of monomer building blocks are incorporated by an enzyme, one might fuse this enzyme with one of the enzymes mentioned above (e.g., the UDP-glykogen synthetase). This would allow the fusion-protein to first incorporate a monomer through reaction of its reactive groups type I, and right thereafter (as the now-incorporated monomer emerges from the active site of the enzyme), the other half of the fusion-protein (e.g., the UDP-glykogen synthetase) would link the functional entity of that monomer to the functional entity of the previous monomer in the complementing template.

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The functional groups (or backbone structures) may have to be protected, in order to not react with the reactive groups or other components of the system during incorpo-

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ration, polymerization and activation. This may be achieved using standard protection groups, some of which are mentioned in (Figure 10).

The polymerization reactions described herein below are divided into two major groups, dependent on whether the funtional entity is held in a fixed oritentation relative to the complementing template.

Group 1: The functional entitles can rotate relative to the complementing elements (and can therefore rotate relative to the complementing template).

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Direct linkage of reactive groups: The reactive group type II of one monomer react directly with the reactive group type II of another monomer.

a). In one example, the functional entity carries two reactive groups X1 and X2 of the same kind. "Same kind" in this respect means that a given X1 can react with both an identical X1 and a non-identical X2. In (Figure 11) X1 and X2 are identical, wherefore they are both symbolized with an X. X may react with another X to form XX (Figure 11). As an example, X might be a thiol (-SH) and the resulting product a disulfide (-SS-). As another example, X could be a coumarin moiety which upon photo-Induction reacts with a coumarin moiety of a neighbouring monomer (Figure

In most cases, the reaction of X with X results in the loss of an atom or a molecular molety; in the case of the thiol, for example, two protons are lost upon disulfide for-

example 1).

25 mation. The fact that XX (the result of the reaction between two reactive groups type II) does not contain all the components of X plus X, Is indicated in (Figure 5, A) where in fact both types of reactive groups (both type I and II) upon reaction forms a molecular entity that is slightly different from the reactive groups (symbolized by overlapping circles in the figure).

b). The two reactive groups type II may be of a different kind. "Different kind" here means that they react with different types of molecules. For example, X and Y might be nucleophiles and electrophiles, respectively. X and Y react to form XY (Figure 12). For longer templated molecules, free rotation of the functional entities relative

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to the complementing template represents a potential problem, if the functional entities do not react until many monomers have been incorporated. In this case, cluster formation (Figure 13) may result, which decreases the amount of full-length, templated polymers. The problem is, however, only significant for longer polymers; from experience with biological display of α-peptides, such as phage-display and polysome-display, it is known that display efficiencies as low as 1 % is enough to isolate peptides with high binding affinity for a given target.

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In certain cases the incorporated monomers react right after their incorporation into the complementing template (at which time the next monomer in the complementing template has not been incorporated yet). Therefore, the last incorporated monomer will react with the second-last incorporated monomer, which is already part of the complementing template. As a result, cluster formation will not be a significant problem in this case.

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X and Y might be an amine and a carboxylic acid. In the presence of carbodlimide, X and Y will react to form an amide XY.

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Another version of this type of polymenization involves the simultaneous polymerization and activation of the polymer (Figure 14). The monomers do not contain a separate linker molety; rather, the polymerization reaction leads to activation (release of the funtional entity from the complementing template). In this scheme, each monomer is incorporated and reacts with the previously incorporated monomer, leading to the previously incorporated monomers release from the complementing template, before the next monomer is incorporated. (Figure 14, example 1) shows the use of this principle for the formation of polyamides, in this case β-peptides. The method may obviously be used for other peptides also, as well as any kind of polyamides.

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By appropriate design of the monomers, one may generate other types of polymer bonds by nucleophilic substitution reactions, including amide, ester, carbamate, carbonate, phosphonate, phosphodiester, sulfonamide, urea, carbopeptide, glycopetide, saccharide, hydrazide, disulfide and peptoid bonds.

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In (Figure 14, example 2) the same principle is applied to a different type of reaction, a "rolling circle polymerization reaction". An alkyl sulfonate is here used as an efficient leaving group, to drive the formation of a secondary amine. The result is a functionalized polyamine, attached at one end to the template that directed its synthesis. In an analogous way, one may generate polyether and poly-thloether using

Iunctionalized polyamine, attached at one end to the template that directed its synthesis. In an analogous way, one may generate polyether and poly-thloether using similar molecular designs. Polymers that can be generated by the use of the principles described in (Figure 14 and 14, example 1 and 2) include oligodeoxynucleotides, oligoribonucleotides, chimeric oligonucleotides, oligonucleotide analogs (e.g., PNA, LNA), peptolds, polypeptides and β-peptides.

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"Fill-In" polymerization: An additional molecule mediates linkage between reactive groups type II from neighbouring monomers.

a). The functional entity carry one or two reactive groups X1 and X2 of the same kind, where X1 cannot react with another X1 or X2. For example, X1 and X2 could be a primary and secondary amine, respectively. In order to polymerize, a compound of the kind Y1-linker-Y2 is added, where Y1 and Y2 are of the same kind. Y can react with X, but is sterically or chemically excluded from reaction with another Y. As a result, a X-Y-Y-X is formed (Figure 15). As an example, X could be an amine, and Y a activated ester. Upon reaction, this would form an ester-ester bond

(X-Y-Y-X) between two functional entities.
It is preferred that the two X of one monomer does not to any significant extent react

with the same Y-linker-Y molecule. This can be prevented e.g. by imposing steric constraints on the molecules, e.g., Ys in the Y-linker-Y molecule are further apart than the Xs in the monomer.

(Figure 15, example 2) provides two examples of "fill-in" polymerization of polyamides. In (Figure 15, example 2, A and B), the reactive groups type II are amines, and the Y-linker-Y molecule is a dicarboxylic acid or an activated di-ester. In either case, the resulting product is a di-amide polymer. Obviously, the kind of X and Y could be switched, so that in the examples X was a carboxylic acid and Y an amine. Other combinations of X and Y, and their resulting bonds, are given in Figure 25,

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which summarizes some of the kinds of polymers that may be generated by the 35 various polymerization principles described in the present patent.

An example is shown in (Figure 15, example 6), where an activated phosphodiester makes up the only reactive group type II of the monomer. Upon reaction with a di-The functional entity may in certain cases contain only one reactive group type II. hydroxy, a phosphodiesterbackbone is formed.

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As yet another example of fill-in polymerization, (Figure 15, example 7) shows the pericyclic reaction of dienes (functional entity) reacting with alkenes (linking molecule), to form a polycyclic compound.

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A general consideration when using the fill-in polymerization principle, is the number is connected to the complementing template through a chiral carbon. The functional ample 4, A), the functional entity contains two primary amines. The functional entity entity may rotate freely around the bond that connects this chiral atom with the comof stereoisomers templated by the same template. For example, in (Figure 15, exmolecules (activated carbonyls, (Y)) will result in the formation of 2^n different isoplementing template. Therefore, the reaction of the amines (X) with the linking mers, where n is the number of residues of the polymer.

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meric polymer, the chirality represents a 1024-fold increase in diversity. This may in certain cases be an advantage, for example if the monomer diversity is low, or if the able. One may then choose to connect the functional entities to the complementing gency of the selection process. Therefore, in certain cases scrambling is not desirdesire is to make short polymers. However, such "scrambling" of the genetic code (i.e., one template encodes different polymer structures) also decreases the strin-The isomers represent a significant increase in diversity. For example, for a 10-

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elements via non-chiral atoms. In (Figure 15, example 4, B) is shown an example of where a complementing element specifies slightly different or entirely different funcfies different isomers (as described above), and scrambling may also involve cases template. Scrambling may involve cases where one complementing element specian achiral atom (nitrogen) connecting the functional entity with the complementing

(Figure 16). X does not react with X or S, and vice cersa. Before, during or after b). The functional entity carries two different reactive groups of type II, X and S

tional entities.

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- ween the functional entities. It is important to ensure that X and S of one functional incorporation of monomers, molecules of the form T-linker-Y are added. X may reantity cannot react with T and Y of one linking molecule. This may be ensured by appropriate design of the structure of the functional entities and linking molecule. act with Y, and S may react with T, leading to formation of X-S-T-Y linkages be-9
- (Figure 16, example 1) provides an example of a functional entity with different reacmolecule with different reactive groups, in this case a phosphine and a ketone (Tive groups type II, in this case an azide and a hydrazide (X and S), and a linking 5
- cules right after their incorporation, or in the case of enzyme-mediated incorporation, For longer templated molecules, free rotation of the functional entities relative to the not react until many monomers have been incorporated. In this case, cluster formacomplementing template represents a potential problem, if the functional entities do plained above. If the linking molecules are present during incorporation of the comolementing elements, the incorporated monomers may react with the linking moletion (Figure 13) may result, which decreases the amount of full-length, templated polymers. The problem is, however, only significant for longer polymers, as ex-ಜ 22
- Zipping" polymerization: The polymerization reaction travels from one end of the emplate to the other.

as soon as they emerge from the active site of the enzyme. Cluster formation will

not be a significant problem in these cases.

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In this approach, the polymertzation reaction is directional, i.e., the reaction cascade starts at one end of the complementing template, and the reactions migrate to the other end of the complementing template, thereby forming a templated polymer.

5 a) General principle (Figure 18). After Incorporation of some or all of the monomer building blocks, polymerization is initiated from one end of the template, and travels down the template. For example, the initiator may be coupled to the first or last complementing element to be incorporated, or it may be coupled to the primer used in DNA polymerase-mediated incorporation of nucleotide-derivatives. Either way,

the initiator will react with the neighbouring monomer's reactive group type II, which induces a change in the functional entity of that monomer, allowing this monomer to react with the next monomer in the chain, and so on. Eventually, all the monomers have reacted, and a polymer has been formed.

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It may be desirable to protect the initiator, keeping it from reacting with the neighbouring monomer until incorporation is complete, whereafter the initiator is deprotected. This allows the experimenter to remove all non-incorporated initiators and complementing elements before activating the initiator, which eliminates reaction in solution between the initiator and the complementing elements.

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Deprotection of the initiator may be by change in pH or temperature, exposure to electromagnetic radiation, or addition of an agent (that removes a protection group, or introduces an initiator at a specific position, or ligates or coordinates to the native initiator, to make it a more potent initiator). The agent could be a chemical catalyst or an enzyme, for example an esterase or peptidase.

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b). Zipping by radical polymerization (Figure 18, example 1). The initiator is a alkyliodide, and the funtional entities contain a double bond. Upon addition of a radical initiator, for example ammoniumpersulfate, AIBN (azobis-isobutyronitrile) or other radical chain reaction initiators, a radical chain reaction is initiated, whereby the alkenes react to form an extended, functionalized alkane. Eventually, the polymer has been made, and it is activated (cleaved from the complementing template, except at one point). The radical remaining at the end of the polymer may be quenched by a radical termination reaction.

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c). Zipping by cationic polymertzation (Figure 18, example 2). The Initiator is a Lewis acid. Upon deprotection with acid or other initiation reagent, a cation is generated. The carbocation attacks the double bond of the neighbouring monomer, and as a result a carbocation is generated in this monomer. Eventually, the full-length

polymer has been formed, and the polymer is activated.

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d). Zipping by nucleophilic (anionic) polymerization (Figure 18, example 3). In this example, the initiator is a protected hydroxyl anion. The functional entity carries a peroxide. Upon deprotection of the initiator the hydroxyl-anion is formed (e.g., by

alkaline deprotection). Under basic conditions, the initiator attacks the neighbouring epoxide at the least hindered carbon in the ring. This in turn generates a hydroxylanion, which attacks the neighbouring epoxide. Eventually, the full polymer is formed, and the polyether may be activated. In this example, all of the linkers that connect the polyether to the complementing template are cleaved.

15 This type of polymerization is also an example of ring-opening polymerization.

 e). Zipping polymerization by ring opening (Figure 19). The general principle of ring-opening polymerization is shown. The initiator attacks the reactive group X of the neighbouring monomer. X is part of a ring structure, and as a result of the reaction between the initiator and X, the ring opens, whereby the other reactive group of

40 ton between the Initiator and X, the ring opens, whereby the other reactive group of the monomer is activated for attack on the next monomer in the array. Polymertzation travels down the strand, and eventually the full-length polymer has been formed. 9-peptide formation by ring-opening polymerization of carboxyanhydrides (Figure 19, example 1). The deprotected initiator, a nucleophilic amine, attacks the most electrophilic carbonyl of the N-thiocarboxyanhydride, to form an amide: CSO is released, generating a primary amine, which then attacks the next monomer in the array. Eventually polymerization is complete, and the polymer may be activated, creating a p-peptide attached to the complementing template or template through its

creating a β-peptide attached to the complementing template or template through its C-terminal end. The principle may be used to form other types of peptides, for example D- and L- form mono- and disubstituted α-peptides, β-peptides, r-peptides, carbopeptides and peptolds (poly N-substituted glycin), and other types of polyamides. Also, the principle can be employed for the generation of other polymers, such as polyesters, polyureas, and polycarbamates.

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g). B-peptide formation by ring-opening polymerization of thiazinanone units (Figure amide. As a result, the ring breaks down to release a free thicketone. This generates an amine, which may now attack the thioester of the next monomer in the array. When polymerization has travelled to the other end of the template, it is activated, generating a β-peptide attached to its template through the C-terminal end. 19, example 2). The deprotected initiator attacks the cyclic thioester, to form an

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form mono- and disubstituted α -peptides, β -peptides, γ -peptides, carbopeptides and peptoids (poly N-substituted glycin), and other types of polyamides. Also, the prin-The principle may be used to form other types of peptides, for example D- and Lciple can be employed for the generation of other polymers, such as polyesters, polyureas, and polycarbamates.

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the initiator leads to a rearrangement of the monomer, which results in activation of h) Zipping polymerization by rearrangement (Figure 20). Upon activation of the inifrom the complementing element. In the attacking monomer, the reaction of Y with X, the other reactive group type II of the monomer (for example, the reorganization phile. Eventually, full-length polymer has been formed, attached at one end to the liator, which in this case could be an electrophile, the reactive group type II of the neighbouring monomer attacks the initiator, and as a result, releases the initiator creates a nucleophile). Then, the next monomer in the array attacks this nucleotemplate that directed its synthesis. 5 20

 Zipping and activation in one step (Figure 21). By appropriate design of the functional entities used for ring-opening polymerization, activation may be achieved as a plementing elements through one of the phenyl groups would lead to activation as a the final polymer to the complementing template, saves the experimenter an activation step (compare Figure 21 and Figure 19). As a specific example, attachment of upside-down, i.e., attach the portion of the ring that does not get incorporated into direct result of the polymentation reaction. By simply turning the functional entity the 2,2-diphenylthiazinanone ring structure of (Figure 19, example 2) to the comresult of the polymerization reaction.

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Group 2: The functional entity cannot rotate freely relative to the complementing element In this embodiment, the X and Y reactive groups type II are held in the desired orientation relative to the complementing template (Figure 22, A). X and Y can therefore react, or react with a linker molecule, without the risk of cluster formation (compare with Figure 13).

bonds to different atoms in the complementing element. (Figure 22, B) provides an tains a dipeptide, and the reactive groups are the amine and the ester moleties, reexample, where the functional entity is covalently coupled to the two bases of a dinucleotide (the complementing element is a dinucleotide, the functional entity con-The functional entity may be held in the fixed orientation by a double bond, or by spectively). 9

Polymers that can be made by this method include all of the polymers mentioned in the non-zipping polymerizations above, for example peptides, amides, esters, carbamates, oximes, phosphodiesters, secondary amines, ethers, etc.

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The figures 23 to 25 relates to how a cluster formation can be avoided by covalent constrains. 20

potential free rotation around the linker-nucleotide bond. A bifunctional FE bears two ure 13 bottom'). This unfavourable situation can be avoided by using fixed functional where 'X' on one FE is meant to react with 'Y' on the neighbour FE either directly or product of say 5 units will be formed ('Figure 13 top'). However, rotation around the groups are arranged so that reaction can take place in two different directions ('Figlinker-bond of some, but not all, linker-FE entities so that the relative orientation of through a cross-linking agent. If all linker-FE units orient Identically with respect to the two functionalities reverses leads to a clustering situation, where the reacting A special situation arises employing bifunctional functional entities (FE) due to a entities thereby preventing rotation around the nucleotide-linker bond. Fixing the different reactive groups 'X' and 'Y', e.g. both a nucleophile and an electrophile, the parent nucleotide, directional polymerization will take place and a complete

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FEs may be obtained by attaching the FE not by one but by two covalent bonds (i.e.

reaction, whereas the additional bond in the latter should be constructed so that also the functionalities, or the two reactive groups may be attached separate 'arms' on a two finkers) to the nucleotide. The additional bond may be formed directly by one of fixed backbone. In the first situation the additional bond may be broken during the this bond is cleavable after reaction, to release the final product.

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position C6 in T/U and C or position C8 or N6 in (deaza)A or (deaza)G, or it could be The primary attachment points of the linker-FE units are typically within the bases of the base or in the sugar part, it could be a second atom in the same base, preferably ond bond should preferably be somewhat distant from this attachment point. That is, deaza G. In order to construct an efficient inhibition of linker-bond rotation, the secthe nucleotides, preferably position C5 in T/U and C or position C7 in deaza A and the second attachment could be anywhere in a neighbour nucleotide, preferably in an atom of the sugar molety, preferably position C2 or C3. Explicit examples are given in Figure 23.

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may be necessary to incorporate by other means than using a polymerase. An al-It should be noted that nucleotides bearing some of these doubly-attached linkers ternative to polymerase incorporation is the imidazole approach described elsewhere herein.

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thereby the most probable product. Therefore, the conformational space covered by amples shown in Figure 23A and 23B. The purpose is to analyse various modes of the linker-FE unit and the zones occupied by the reactive groups needs to be estipolymerization a series of computer calculations have been performed on two exattack for each linker-FE construction, estimating the most probable reaction and In order to show the effect of covalently constraining the FE to ensure directional mated.

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different searching methods and is standard knowledge within the field. For systems of the size mentioned in this text it is not possible to perform a converged conformabe performed employing various different software and within these programs using can be estimated by doing a conformational search. Conformational searches can The conformational space of a specific linker-FE system, i.e. the range of the FE, complete potential energy surface has been covered and thereby that the located ional search, that is, to ensure that enough steps have been taking so that the

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tion distance. Efficient conformational searching methods employing a rather limited ever, the purpose of these calculations is to get a picture of the space allowed to be covered by a linker-FE unit and thereby to estimate the most likely approach of attack between two FEs and the possibility for the reacting groups to get within reacminimum energy conformation is truly the global minimum for the molecule. Hownumber of steps fulfil this purpose.

different overall configurations, by which is meant an overall arrangement of the two corresponds to one specific configuration, and four linker-FE units arranged e.g. with ple rotation about single bonds. Different conformations may in addition give rise to reactive groups on all modified nucleotides that give rise to one specific direction of conformations are possible, but all of these result in the same 'most probable' prod-By conformations are here meant Individual structural orientations differing by simtwo 'X's' pointing in one direction and the two other in the opposite direction corresponds to another specific configuration. Within one configuration many different reaction. That is, four linker-FE units arranged with all 'X's' in the same direction act since the overall orientation (direction) of reactive groups is preserved. 5 우

The calculations performed in this investigation have been performed employing the backage a series of different searching protocols are available, including the 'Mixed Monte Carlo Multiple Minimum/Low Mode' method (MCMM/LM), shown to be very MacroModel7.2 software from Schrödinger Inc (MMOD72). Within this program effective in locating energy minima for large complicated systems.

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Computational details 25

Double-stranded DNA with the base sequence 5'-GCTTTTTAG-3' (upper strand) (example displayed in Figure 24) or 5'-GCTTTTAG-3' (upper strand) (example displayed in Figure 25) was built using HyperChem7 from HyperCube Inc in the most requent B-conformation. The linker-FE units were built using ChemDraw Ultra 6.0 nto MMOD72. The linkers were then fused to the corresponding nucleotides using calcutations all DNA atoms were kept frozen, that is, were not allowed to move, in and Chem3D Ultra 6.0 from ChemOffice. Linker-FE units and DNA were imported the build feature in MMOD72, fusing the methyl carbon atom of the T nucleotides with the appropriate linker atom, in effect creating a modified U nucleotide. In all order to decrease the size of the systems and to avoid distortions within the DNA

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of between 100 and 1000 applied. Without the constraint none of the MMOD72 force N1, C6, C5 and the first linker atom was set to 180.0 degrees and a force constant employing the OPLS_AA force field supplied in MMOD72, It was necessary to constrain the dihedral angle bridging the nucleotide and the linker, i.e. the dihedral of fields preserved the plane, presumably due to a too weak out-of-plane force constrand. The total system was energy minimised (keeping the DNA atoms frozen) stant for this particular dihedral.

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distance travelled by the fastest moving atom of 3 and 8 A, respectively. Depending 0.05 kJ/mol). The chirality of chiral atoms was preserved during the calculations. In each conformer was minimized by 500-1000 PR Conjugate Gradient steps (this re-2000 steps with an energy cutoff of 50 kJ/mol, and with a minimum and maximum on the specific size of the system, 11-19 torsions were allowed to vary, and finally addition, for the systems with covalently constrained linkers one ring closure bond (either the formed amide bond or the base-S bond) was chosen within each ring. sulted in most conformers being minimized to within a convergence threshold of Conformational analyses were performed using the MCMM/LM method, running

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Employing the same linker-FE unit without the second attachment point corresponds linked to the 3' base. When reaction takes place the amino group from one dinucleo-Linker 1A is constructed from a β -dipeptide, with the amino end connected to the 5' base via a disulfide bond prone for reductive cleavage and the carboxy-end directly FE bears two reactive groups on separate arms and has free rotation about the nu-FE Is increased. Using this approach, a series of valid attachment points exist; Figtide-linker-FE unit will break the ester bond of the preceding dinucleotide-linker-FE. to employing dipeptides on one-nucleotide-spaced mononucleotides. Such a linkerties via a breakable bond to a nelghbour nucleotide. In effect, this means that dinucleotides in stead of mononucleotides are employed and also that the length of the One way of creating a second attachment point is to link one of the functional enti-Running 2000 conformational search steps of the singly-attached linker results in ure 23A is an example of attachment to the same position of the neighbour base. cleotide-linker bond and is therefore an example of a bifunctional linker-FE which bears the risk of cluster formation in case of lack of directional polymerization.

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490 unique conformations (849 conformations after 500 mlnImIzation steps, 490

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energy conformation which results In a complete product has rank 8 and is shown in groups upwards and all carboxy groups downwards, and the two reactions that are Figure 24A and the resulting most probable product (still attached the DNA backbone) in Figure 24B. As can be seen, the reactive groups arrange with all amino after additional 500 steps) with the 'global' minimum located once. The lowest-

- product is shown as Figure 24H. However, by far the most conformations, including ormed an amide bond, but the 5' linker-FE unit has the opposite overall orientation the 'global' minimum, do not have this overall configuration. Figure 24C shows the Figure 24D. As can be seen, this arrangement of the reactive groups results in the conformation of second highest rank and the most probable product is depicted in resulting in two carboxy groups (one from the 5' linker-FE unit and one from the required to give a complete product are straight-forward. The released complete formation of an incomplete product. The two linker-FE units in the 3' end have
- the dimer is depicted in Figure 22. Of the 364 unique conformations within 10 kJ/mol of the located minimum approximately 330 results in the formation of various incommerged 3' linker-FE units) being the two close reactive groups and thus no reaction is possible. Release of this product therefore results in a dimer (and a monomer); plete products. 0 5
- the DNA backbone). Release of this product gives the compiete three-unit product, Running 2000 conformational search steps of the doubly-attached linker results In 125 unique conformations with the 'global' minimum located nine times. This minilinker, giving rise to only one probable product shown in Figure 24F (still attached Clearly, this overall configuration is the only one possible for the doubly-attached mum energy conformation is shown In Figure 24E, where all FEs are seen to arrange with the amino groups pointing downwards and carboxy groups upwards. 8 33
- important issue is the difference in complete products formed. The FEs employed in result in (many) configurations unable to form complete products. However, another Thus, this example shows first of all that rotation around nucleotide-linker bonds do this example are constructed from unsubstituted β-amino acids and therefore there ever, using singly-attached FEs the polarity of the formed products can change (i.e. Is no difference between the complete products shown in Figure 24 G and H. Howfree amino group from the 3' attached FE or free amino group from the 5' attached ဓ 33

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FE) and thereby potentially very different products can be formed. By employing fixed functional entities only one overall configuration is possible and only one product with one specific polarity can be formed.

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Another possibility of attachment point is the sugar of the parent nucleotide, as exemplified in Figure 23 B, C, and D. This choice allows the employment of mononucleotides in stead of linked dinucleotides as mentioned above. Both hydrogens of C2 can be replaced by linker atoms, however, for shorter ring structures it is preferred to use the one facing the same plane as the base does. Carbon 3 of the sugar molety forms a linkage to the phosphate group, but there is still one attachment possibility left which can be utilised for linker fixation purposes. The same holds for C1, however the space around this substitution possibility is limited. Carbon atoms 4 and 5 of the sugar moiety are quite distant from the base attachment point and therefore require large ring systems to be utilised for this purpose.

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Linker-FE 18 is constructed from a y-amino acid attached via a disulfide bond prone for reductive cleavage to the C5 position of T/U. The linker-FE unit 18 is therefore another typical example of a bifunctional linker-FE system capable of rotation of the nucleotide-linker bond which bears the risk of cluster formation due to lack of directional polymerization. A fixation of this FE is shown in example 18 (right) and utilises the C2 position at the same side of the plane as the base. The FE now contains a y-amino acid linked through the carboxyl group to the sugar via a hydrolysable ester bond and in the amino end to the C5 position of T/U via a disulfide bond prone for oxidation. Doubly-attached linker-FE unit 18 is therefore a bifunctional linker-FE with one reactive group free and the other providing the second attachment. Using almost the same linker-FE unit but letting the carboxyl end free by introducing a second ester group as a hydrolysable linker is shown in example 10. Computational analyses of linker 18.

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Conformational searches of the two different schemes clearly reveal the effect of preventing rotation of the linker bond by additional covalent attachment.

Running 2000 conformational search steps of the singly-attached linker-FE results in 445 unique conformations with the 'global' minimum located once. This conformation is shown in Figure 25A and the resulting most probable product (still attached the DNA backbone) in Figure 25B. As can be seen this product is the complete

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product, that is, all four units are linked together via amide bonds. The released complete product is shown as Figure 25G. However, many other overall configurations are possible for this system, with one example shown in Figure 25C. The most probable product resulting from the 3C configuration is depicted in Figure 25D and as can be seen, this arrangement of reactive groups results in the formation of an incomplete product, that is, the linker-FE units are linked two and two together with no possibilities of a merging reaction. Release of this product results in two dimers, depicted in Figure 25H. Of the 334 unique conformations within 10 kJ/mol of the located minimum approximately 215 results in formation of various incomplete prod-

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Running 2000 conformational search steps of the doubly-attached FE results in 386 unique conformations with the 'global' minimum located twice. This conformation is shown in Figure 25E and the resulting most probable product (still attached the DNA backbone) in Figure 25F. However, since there are no possibilities of interchange of reactive groups, the conformations differ only by minor variations in dihedrals (e.g. rotation of the CH₂NH₂ group). Clearly, only one overall configuration is possible for the doubly-attached FE, giving rise to only one probable product, the complete four-

unit product (Figure 25G).

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Thus, the computational investigations clearly show that there is extensive rotation around nucleotide-linker bonds and that this flexibility will result in a significant proportion of the formed products not being complete. The calculations also show that using covalently fixed functional entities is one way to prevent linker rotation and thereby effectively secure unidirectional polymerisation. In addition, the complete products that do result from using unconstrained FEs form a diverse group, since there is more than one possibility of arranging the reactive groups in a way that allows reactions between all units to happen. Naturally, these tendencies will be even more pronounced using more than three to four linker-FE units as was applied in these examples.

Building blocks capable of transferring functional entitles.

The following section describes the formation and use of monomer building blocks 35 capable of transferring a functional entity from one monomer building block to an-

other monomer building block, i.e. two functional entities of two monomer building blocks react, whereby one functional entity is cleaved from its monomer building block under the conditions applied.

General Section

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Protection and deprotection of maleimide derivatives:

Malelmide derivatives (e.g. R = H, alkyl, aryl, aryl, alkoxy etc.) may at any step below, be present in a protected form. Protection is achieved by reaction with furan. Deprotection may be achieved by thermolysis, as described by Masayasu et al., J. Chem. Soc., Perkin Trans. 1 (1980) 2122.

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A. Acylation reactions

15 General route to the formation of acylating monomer building blocks and the use of

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N-hydroxymaleimide (1) may be acylated by the use of an acylchloride e.g. acetylchloride or alternatively acylated in e.g. THF by the use of dicyclohexylcarbodiimide or disopropylcarbodiimide and acid e.g. acetic acid. The intermediate may be subjected to Michael addition by the use of excess 1,3-propanedithiol, followed by reaction with either 4,4-dipyridyl disulfide or 2,2'-dipyridyl disulfide. This intermediate (3) may then be loaded onto an oligonucleotide carrying a thiol handle to generate the monomer building block (4). The reaction of this monomer building block with an arnine carrying monomer building block is conducted as follows:

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The template oligonucleotide (1 nmol) Is mixed with a thio oligonucleotide loaded with a building block e.g. (4) (1 nmol) and an amino-oligonucleotide (1 nmol) in hepes-buffer (20 µL of a 100 mM hepes and 1 M NaCl solution, pH=7.5) and water (39 uL). The oligonucleotides are annealed to the template by heating to 50 °C and cooled (2 °C/ second) to 30 °C. The mixture is then left o/n at a fluctuating temperature (10 °C for 1 second then 35 °C for 1 second), to yield template bound (5).

B. Alkylation and C. Vinylation reactions

General route to the formation of alkylating/vinylating monomer building blocks and

Alkylating monomer building blocks may have the following general structure: ß

R¹ = H, Me, Et, iPr, Cl, NO₂ R² = H, Me, Et, iPr, Cl, NO₂

R1 and R2 may be used to tune the reactivity of the sulphate to allow appropriate crease reactivity. Ortho substituents to the sulphate will due to steric reasons direct reactivity. Chloro and nitro substitution will increase reactivity. Alkyl groups will deincoming nucleophiles to attack the R-group selectively and avoid attack on sulphur.

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₩.¥ <u>9</u> Ξ 3-Aminophenol (6) is treated with malelc anhydride, followed by treatment with an acid e.g. H₂SO₄ or P₂O₅ and heat to yield the malefmide (7). The ring closure to the maleimide may also be achieved when an acid stable O-protection group is used by treatment with or Ac₂O with or without heating, followed by O-deprotection. Alternatively reflux in Ac₂O, followed by O-deacetylation in hot water/dioxane to yield (7).

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Further treatment of (7) with SO₂Cl₂ with or without triethylamine or potassium carwhich may be isolated or directly further transformed into the aryl alkyl sulphate by the quench with the appropriate alcohol, in this case MeOH, whereby (9) will be formed. The organic building block (9) may be connected to an oligo nucleotide, as bonate in dichloromethane or a higher boiling solvent will yield the intermediate (8),

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The reaction of the alkylating monomer building block (10) with an amine carrying monomer building block may be conducted as follows:

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°C/ second) to 30 °C. The mixture is then left o/n at a fluctuating temperature (10 °C The oligonucleotides are annealed to the template by heating to 50 °C and cooled (2 The template oligonucleotide (1 nmol) is mixed with a thio oligonucleotide loaded buffer (20 μL of a 100 mM hepes and 1 M NaCl solution, pH=7.5) and water (39 uL). for 1 second then 35 °C for 1 second), to yield the template bound methylamine with a building block (1 nmol) (10) and an amino-oligonucleotide (1 nmol) in hepes-

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the chlorosulphonate (8 above) with an alcohol, the intermediate chlorosulphate is scribed above for an alkylating monomer building block. Although instead of reacting A vinylating monomer building block may be prepared and used similarily as deisolated and treated with an enclate or O-trialkyIsIIyIenolate with or without the presence of fluoride. E.g.

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Formation of the vinylating monomer building block (13):

The thiol carrying oligonucleotide in buffer 50 mM MOPS or hepes or phosphate pH 7.5 is treated with a 1-100 mM solution and preferably 7.5 mM solution of the organic building block (12) in DMSO or alternatively DMF, such that the DMSO/DMF concentration is 5-50%, and preferably 10%. The mixture is left for 1-16 h and preferably 2-4 h at 25 °C. To give the vinylating monomer building block (13).

ing block to give an enamine (14a and/or 14b) or e.g. react with an carbanion to The sulfonylenolate (13) may be used to react with amine carrying monomer bulldyield (15a and/or 15b). E.g.

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The reaction of the vinylating monomer building block (13) and an amine or nitroalkyl carrying monomer building block may be conducted as follows:

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kyl-oligonucleotide (1 nmol) in 0.1 M TAPS, phosphate or hepes-buffer and 300 mM nealed to the template by heating to 50 °C and cooled (2 °C/ second) to 30 °C. The with a building block (1 nmol) (13) and an amino-oligonucleotide (1 nmol) or nitroalfor 1 second), to yield template bound (14a/b or 15a/b). Alternative to the alkyl and mixture is then left ofn at a fluctuating temperature (10 °C for 1 second then 35 °C The temptate oligonucleotide (1 nmol) is mixed with a thio oligonucleotide loaded NaCl solution, pH=7.5-8.5 and preferably pH=8.5. The oligonucleotides are an-

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(however with R" instead as alkyl or vinyl), described below, be prepared from (28; with the phenyl group substituted by an alkyl group) and (29), and be used as alkyating and vinylating agents. 5

vinyl sulphates described above may equally effective sulphonates as e.g. (31)

D. Alkenylidation reactions ឧ

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General route to the formation of Wittig and HWE monomer building blocks and use

Commercially available building block (16) may be transformed into the NHS ester

(17) by standard means, i.e. DCC or DIC couplings. G

17) be P-alkylated with an alkylhalide and then be coupled onto an amine carrying An amine carrying oligonucleotide in buffer 50 mM MOPS or hepes or phosphate pH centration is 5-50%, and preferably 10%. The mixture is left for 1-16 h and preferably 2-4 h at 25 °C. To give the phosphine bound monomer building block (18). This halide, e.g. N,N-dimethyl-2-lodoacetamide as a 1-100 mM and preferably 7.5 mM solution in DMSO or DMF such that the DMSO/DMF concentration is 5-50%, and preferably 10%. The mixture is left for 1-16 h and preferably 2-4 h at 25 °C. To give the monomer building block (19). Alternative to this, may the organic building block 7.5 is treated with a 1-100 mM solution and preferably 7.5 mM solution of the organic building block in DMSO or alternatively DMF, such that the DMSO/DMF conmonomer building block is further transformed by addition of the appropriate alkyl-

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An aldehyde bound monomer building block (20), e.g. formed by the reaction between the NHS ester of 4-formylbenzoic acid and an amine carrying oligonucleotide, oligonucleotide to yield (19).

using conditions similar to those described above, will react with (19) under slightly

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alkaline conditions to yield the alkene (21).

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The reaction of monomer building blocks (19) and (20) may be conducted as fol-

The template oligonucleotide (1 nmol) is mixed with monomer building block (19) (1 nmol) and (20) (1 nmol) in 0.1 M TAPS, phosphate or hepes-buffer and 1 M NaCl solution, pH=7.5-8.5 and preferably pH=8.0. The reaction mixture is left at 35-65 °C preferably 58 °C over night to yield template bound (21).

As an atternative to (17) may phosphonates (24) be used instead. They may be prepared by the reaction between diethylchlorophosphite (22) and the appropriate carboxy carrying alcohol. The carboxylic acid is then transformed into the NHS ester (24) and the process and alternatives described above may be applied. Although

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Instead of a simple P-alkylation, the phosphite will undergo Arbuzov's reaction and generate the phosphonate. Monomer building block (25) benefits from the fact that it is more reactive than its phosphonium counterpart (19).

E. Transition metal catalyzed arylation, hetaylation and vinylation reactions

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Electrophilic monomer building blocks (31) capable of transferring an aryl, hetaryl or vinyl functionality may be prepared from organic building blocks (28) and (29) by the use of coupling procedures for malelmide derivatives to SH-carrying oligonucleotide's described above. Alternative to the malelmide, may NHS-ester derivatives prepared from e.g. carboxybenzensulfonic acid derivatives, be used by coupling of such to an amine carrying oligonucleotide. The R-group of (28) and (29) is used to tune the reactivity of the sulphonate to yield the appropriate reactivity.

The transtion metal catalyzed cross coupling is conducted as follows:

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A premix of 1.4 mM Na₂PdCl₄ and 2.8 mM P(p-SO₃C₈H₄)₃ in water left for 15 min was added to a mixture of the template oligonucleotide (1 nmol) and monomer building block (30) and (31) (both 1 nmol) in 0.5 M NaOAc buffer at pH=5 and 75 mM NaCl (final [Pd]=0.3 mM). The mixture is then left of at 35-65 °C preferably 58 °C, to yield template bound (32).

R" = aryl, hetaryl or vinyl

Corresponding nucleophilic monomer building blocks capable of transferring an aryh, hetaryl or vinyl functionality may be prepared from organic building blocks type (35).

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This is available by estrification of a boronic acid by a diol e.g. (33), followed by transformation into the NHS-ester derivative. The NHS-ester derivative may then be coupled to an oligonucleotide, by use of coupling procedures for NHS-ester derivatives to amine carrying oligonucleotide's described above, to generate monomer

5 building block type (37). Alternatively, may materimide derivatives be prepared as described above and loaded onto SH-carrying oligonucleotide's.

The transtion metal catalyzed cross coupling is conducted as follows:

A premix of 1.4 mM Na₂PdCl₄ and 2.8 mM P(p-SO₃C₆H₄)₈ in water left for 15 min was added to a mixture of the template oligonucleotide (1 nmol) and monomer building block (36) and (37) (both 1 nmol) in 0.5 M NaOAc buffer at pH=5 and 75 mM NaCl (final [Pd]=0.3 mM). The mixture is then left o/n at 35-65 °C preferably 58 °C, to yield template bound (38).

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(36)

R = aryl, hetaryl or vinyl

F. Reactions of enamine and enolether monomer building blocks

Monomer building blocks loaded with enamines and enolethers may be prepared as c)

For Z=NHR (R=H, alkyl, aryl, hetaryl), a 2-mercaptoethylamine may be reacted with densed to a ketone or an aldehyde under dehydrating conditions to yield the a dipyridyl disulfide to generate the activated disulfide (40), which may then be conenamine (41).

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For Z=OH, 2-mercaptoethanol is reacted with a dipyridyl disulfide, followed by Otosylation (Z=OTs). The tosylate (40) may then be reacted directly with an enolate or The enamine or enolate (41) may then be coupled onto an SH-carrying oligonucleoin the presence of fluoride with a Otrialkylsilylenolate to generate the enolate (41).

tide as described above to give the monomer building block (42).

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$$(39) \qquad (40) \qquad (41)$$

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deotide like (44) or alternatively an alkylhalide carrying oligonucleotide like (43) as The monomer building blocks (42) may be reacted with a carbonyl carrying oligonu-

mM NaCl solution, pH=7.5-8.5 and preferably pH=7.5. The reaction mixture is left at 35-65 °C preferably 58 °C over night or alternatively at a fluctuating temperature (10 °C for 1 second then 35 °C for 1 second) to yield template bound (46), where Z=O or NR. For compounds where Z=NR slightly acidic conditions may be applied to The template oligonucleotide (1 nmol) is mixed with monomer building block (42) (1 nmol) and (43) (1 nmol) in 50 mM MOPS, phosphate or hepes-buffer buffer and 250 yield product (46) with Z=O.

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nmol) and (44) (1 nmol) in 0.1 M TAPS, phosphate or hepes-buffer buffer and 300 mM NaCl solution, pH=7.5-8.5 and preferably pH=8.0. The reaction mixture is left at 35-65 °C preferably 58 °C over night or alternatively at a fluctuating temperature (10 °C for 1 second then 35 °C for 1 second) to yield template bound (45), where Z=O or NR. For compounds where Z=NR slightly acidic conditions may be applied to The template oligonucleotide (1 nmol) is mixed with monomer building block (42) (1 yield product (45) with Z=0.

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<u>8</u> £ € Enolethers type (13) may undergo cycloaddition with or without catalysis. Similarly, may dienolethers be prepared and used. E.g. by reaction of (8) with the enclate or trialkylsilylenolate (in the presence of fluoride) of an $\alpha \beta$ -unsaturated ketone or aldehyde to generate (47), which may be loaded onto an SH-carrying oligonucleotide, to yield monomer building block (48).

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The diene (49), the ene (50) and the 1,3-dipole (51) may be formed by simple reac-

sponding organic building block. Reaction of (13) or alternatively (31, R"=vinyl) with dienes as e.g. (49) to yield (52) or e.g. 1,3-dipoles (51) to yield (53) and reaction of (48) or (31, R"=dienyl) with enes as e.g. (50) to yield (54) may be conducted as foltion between an amino carrying oligonucleotide and the NHS-ester of the correlows: 2

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(48) (1 nmol) and (49) or (50) or (51) (1 nmol) in 50 mM MOPS, phosphate or The reaction mixture is left at 35-65 °C preferably 58 °C over night or alternatively at The template oligonucleotide (1 nmol) is mixed with monomer building block (13) or hepes-buffer buffer and 2.8 M NaCl solution, pH=7.5-8.5 and preferably pH=7.5. a fluctuating temperature (10 °C for 1 second then 35 °C for 1 second) to yield tem-

plate bound (52), (53) or (54).

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elements and the functional entity) may be done by changes in pH and/or temperaesterases and nucleases. A list of cleavable linkers and the conditions for cleavage ture, addition of reactants or catalysts, enzymes or ribozymes, or light, UV or other electromagnetic radiation, etc. Particularly relevant enzymes include proteases, Activation (cleavage of some or all of the linkers connecting the complementing is shown in (Figure 10).

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which is cleaved by acid, the 2-[(tert-butyldiphenylsiloxy)methyl]benzoic acid molety acid moiety which provides a linker cleavable by the combination of alkaline phoswhich is celavable with fluoride, and the phosphate of a 2-hydroxymethyl benzoic Other cleavable linkers include the 4-hydroxymethyl phenoxyacetic acid molety, phatase treatment followed by treatment with mild alkaline treatment. 우

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and the functional entities, thereby obtaining a polymer physically linked through just the nature of the linker is not considered an essential feature of this invention. The is in the range from the length of just one bond, to a chain length of about 20 atoms. polymer can vary widely, but for the purposes of the invention, preferably the length to selectively cleave all but one of the linkers between the complementing template one linker to the template that templated its synthesis. This intact linker should af-In most cases, it is desirable to have at least two different types of linkers connectng the complementing elements with the funtional entities. This way, it is possible ect the activities of the attached polymer as little as possible, but other than that, size of the linker in terms of the length between the template and the templated

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Selection and screening of templated molecules

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lytic activity may be performed by affinity selections on transition-state analog affinity polysome-displayed or mRNA-protein fusion displayed peptides. Selection for cata-Selection or screening of the templated molecules with desired activities (for examassay) may be performed according to any standard protocol. For example, affinity selections may be performed according to the principles used for phage displayed, ple binding to particular target, catalytic activity, or a particular effect in an activity

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template

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95(18):10523-8). Screening for a desired characteristic may be performed according columns (Baca et al. , Proc. Natl. Acad. Sci USA. 1997; 94(19);10063-8), or by functionbased selection schemes (Pedersen et al., Proc. Natl. Acad. Sci. USA. 1998, to standard microtiter plate-based assays, or by FACS-sorting assays.

Use of libraries of templated molecules

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binding of the selected molecule should be selective in that they only coordinate to a these binding molecules should be possible to use e.g. as therapeutic agents, or as molecules capable of binding to different targets. The template-displaying molecule specific target and thereby prevent or induce a specific biological effect. Ultimately, The present invention is also directed to approaches that allow selection of small technology contains a built-in function for direct selection and amplification. The Selection of template-displaying molecules that will bind to known targets diagnostic agents.

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molecular structures (e.g. nucleic acids and proteins, inciuding enzymes, receptors, method provides a rapid means for Isolating and Identifying molecule ligands which antibodies, and glycoproteins); signal molecules (e.g. cAMP, inositol triphosphate, peptides, prostagiandins); and surfaces (e.g. metal, plastic, composite, glass, ceselections, or assays to assess the effect of binding of a molecule ligand on the Femplate-displaying molecule libraries can easily be combined with screenings, function of the target. In a more specific embodiment, the template-displaying bind to supra-molecular, macro-supra-molecular, macro-molecular and lowramics, rubber, skin, tissue).

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Specifically, selection or partitioning in this context means any process whereby the target pair, can be separated from template-displaying molecules not bound to the target molecule. Selection can be accomplished by various methods known in the tempiate-displaying molecule complex bound to a target molecule, the complex-

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The selection strategy can be carried out so it allows selection against almost any target. Importantly, no steps in this selection strategy need any detailed structural

called hot-spot regions (Wells, et al. (1993) Recent Prog. Hormone Res. 48; 253-

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molecules in the library to a given target. However, in some applications, if needed, functionality can also be included analogous to selection for catalytic activity using driven by the binding affinity involved in the specific recognition/coordination of the phage display (Soumillion et al. (1994) J. Mol. Biol. 237: 415-22; Pedersen et al. information of the target or the molecules in the libraries. The entire process Is

(1998) PNAS. 18: 10523-10528). Example of various selection procedures are de-

scribed below.

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This built-in template-displaying molecule selection process is well sulted for optimibinding molecules and ends with the optimized binding molecule. The single procedures in each step are possible to automate using various robotic systems. This is zations, where the selection steps are made in series starting with the selection of because there is a sequential flow of events and where each event can be per-9

ormed separately. In a most preferable setting, a sultable tempiate-displaying molefinally generates the optimized binding molecule. Even more preferably, this process should run without any need of external work outside the robotic system during the cule library and the target molecule are supplied to a fully automatic system which antire procedure. 5

sites) on different proteins are more prone to bind molecules that other more neutral plate-displaying molecules. This will be dependent on the precise bInding mode and However, it is known that functional sites (e.g. protein-protein interaction or catalytic smaller region that seems to be primarily responsible for the binding energy, the so tentially coordinate to any known or unknown target. The region of binding on a target could be into a catalytic site of an enzyme, a binding pocket on a receptor (e.g. GPCR), a protein surface area involved in protein-protein interaction (especially a The libraries of template-displayed molecules will contain molecules that could pohot-spot region), and a specific site on DNA (e.g. the major groove). The templatedisplaying molecule technology will primarily identify molecules that coordinate to the particular binding-site the template-displaying molecules occupy on the target. the target molecule. The natural function of the target could either be stimulated surface areas on a protein. In addition, these functional sites normally contain a (agonized) or reduced (antagonized) or be unaffected by the binding of the tem-20

262). This phenomenon will increase the possibility to directly select for small mole cules that will affect the biological function of a certain target

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(Roberts, et al. (1997) Proc. Natl. Acad. Sci. 94: 12297-302). However, the enclosed invention are the amplification of the templates and incorporation and reaction of the monomer building blocks. The amplification and Incorporation and the Incorporation translation process on the ribosome complex. The necessary steps included in this leins (Marks et al. (1992) J. Biol. Chem. 267: 16007-16010) and antibodies (Winter direct selection of target-specific small non-peptide molecules independently of the (1985) Science 228: 1315-1317). Phage display selection has been used success-(Mattheakis et al. (1994) Proc. Natl. Acad. Sci. 91: 9022-9026) and mRNA display invention, the template-displaying molecule technology, will for the first time allow The template-displaying molecule technology of the invention will permit selection fully on peptides (Wells & Lowman. (1992) Curr. Op. Struct. Biol. 2, 597-604) proet al. (1994) Annu. Rev. Immunol. 12: 433-455). Similar selection procedures are procedures analogous to other display methods such as phage display (Smith also exploited for other types of display systems such as ribosome display and reaction are either done in the same step or in a sequential process.

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The Inkage between the templated molecule (displayed molecule) and DNA replicaunknown antigens (epitopes) and use these binding molecules for Identification and binding molecules against any known target. In addition, this technology will also tion unit (coding template) allows a rapid Identification of binding molecules using various selection strategies. This invention allows a broad strategy in Identifying allow discovery of novel unknown targets by isolating binding molecules against validation (see section "Target identification and validation"). ឧ 22

emplate-displayed molecules; Elution of the template-displayed molecules bound to cules. A typical selection procedure against a purified target will include the following major steps; Generation of a template-displaying molecule library: Immobilization of As will be understood, selection of binding molecules from the templata-displaying the target molecule using a sultable immobilization approach; Adding the library to the immobilized target; Amplification of enriched template-displaying molecules for molecule libraries can be performed In any format to identify optimal binding moleallow binding of the temptate-displayed molecules; Removing of the non-binding

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dentification by sequencing or to Input for the next round of selection. The general steps are schematically shown in Figure 39.

displaying molecule library is to use the bio-panning method. In this technique, the in a preferred embodiment, a standard selection protocol using a template-

larget (e.g. protein or peptide conjugate) is immobilized onto a solid support and the hat are selected and enriched. However, the selection procedure requires that the bound template-displayed molecules can be separated from the unbound ones, i.e. emplate-displayed molecules that potentially coordinate to the target are the ones hose in solution. There are many ways in which this might be accomplished as 2

known to ordinary skilled in the art.

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nany orders of amplification, I.e. every single template-displayed molecule enriched The first step in the affinity enrichment cycle (one round as described in Figure 1) Is the stringent washing, is then eluted with, e.g. acid, chaotropic safts, heat, competiwhen the template-displayed molecules showing low affinity for an Immobilized tarattached to the target. The enriched population, remaining bound to the target after tive elution with the known ligand or proteolytic release of the target/femplate molein the first selection round participates in the further rounds of selection at a greatly get are washed away, leaving the strongly binding template-displayed molecules cules. The eluted template-displayed molecules are suitable for PCR, leading to

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assaying the proportion of template-displaying molecules which remain bound to the increased copy number. After typically three to ten rounds of enrichment a populamolecules which bind most strongly to the target. This is followed quantitatively by tion of molecules is obtained which is greatly enriched for the template-displayed mmobilized target. The variant template sequences are then individually sequenced.

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select from the template-displaying molecules, simply by sedimenting the beads in a immobilisation of the target (peptide, protein, DNA or other antigen) on beads might including for instance attachment to -NH2 groups and -SH groups. Magnetic beads be useful where there is doubt that the target will adsorb to the tube (e.g. unfolded bench centrifuge. Alternatively, the beads can be used to make an affinity column argets eluted from SDS-PAGE gels). The derivatised beads can then be used to and the template-displaying libraries suspension recirculated through the column. There are many reactive matrices available for immobilizing the target molecule, are essentially a variant on the above; the target is attached to magnetic beads

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which are then used in the selection. Activated beads are available with attachment

sites for -NH₂ or -COOH groups (which can be used for coupling). The target can be also be blotted onto nitrocellulose or PVDF. When using a blotting strategy, it is important to make sure the strip of blot used is blocked after immobilization of the tar-

5 get (e.g. with BSA or similar protein).

In another preferred embodiment, the selection or partitioning can also be performed using for example: Immunoprecipitation or Indirect Immunoprecipitation were the target molecule is captured together with template-displaying binding molecules; affinity column chromatography were the target is immobilized on a column and the template-displaying libraries are flowed through to capture target-binding molecules; gel-shift (agarose or polyacry/amide) were the selected template-displaying molecules migrate together with the target in the gel; FACS sorting to localize cells that coordinates template-displaying molecules; CsCl gradient centrifugation to isolate the target molecule together template-displaying binding molecules; Mass spectroscopy to Identify target molecules which are labelled with template-displaying molecules; etc., without limitation. In general, any method where the template-displaying molecules not bound to the target is useful.

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20 Table 2: Examples of selection method possible to use to identify binding molecules using the template-displaying technology.

Target Method of choice	eceptors Direct Immobilization, Immunoprecipitation, affinity column, FACS sorting, MS.	e receptor Cell-surface subtraction selection, FACS sorting. Affinity column.	nhibitors Direct immobilization, Immunoprecipitation, affinity column, FACS sorting, MS.	Spitopes Cell-surface subtraction selection, in-vivo
Type of Target	Soluble receptors	Cell surface receptor	Enzyme inhibitors	Surface epitopes

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Elution of template-displayed molecules can be performed in different ways. The binding molecules can be released from the target molecule by denaturation, acid, or chaotropic salts and then transferred to another vial for amplification. Alterna-

tively, the elution can be more specific to reduce the background. Elution can be accomplished using proteolysis to cleave a linker between the target and the immobilizing surface or between the displaying molecule and the template. Also, elution can be accomplished by competition with a known ligand. Alternatively, the PCR reaction can be performed directly in the washed wells at the end of the selection reaction.

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A possible feature of the invention is the fact that the binding molecules need not be elutable from the target to be selectable since only the encoding template DNA is needed for further amplification or cloning, not the binding molecule itself. It is known that some selection procedure can bind the most avid ligands so tightly as to be very difficult to elute. However the method of the invention can successfully be practiced to yield avid ligands, even covalent binding ligands.

Alternative selection protocol includes a known ligand as fragment of each displayed molecule in the library. That known ligand will guide the selection by coordinate to a defined part on the target molecule and focus the selection to molecules that binds to the same region. This could be especially useful for increasing the affinity for a ligand with a desired biological function but with a too low potency.

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A further aspect of the present invention relates to methods of increasing the diversity or complexity of a single or a mixture of selected binding molecules. After the initial selection, the enriched molecules can be altered to further increase the

- chemical diversity or complexity of the displayed molecules. This can be performed using various methods known to the art. For example, using synthesized random-ized oligonucleotides, spiked oligonucleotides or random mutagenesis. The randomization can be focused to allow preferable codons or localized to a predetermined portion or sub-sequence of the template nucleotide sequence. Other preferes able method is to recombine templates coding for the binding molecules in a similar
- able method is to recombine templates coding for the binding molecules in a similar manner as DNA shuffling is used on homologous genes for proteins (Stemmer (1994) Nature 370:389-91). This approach can be used to recombine initial libraries or more preferably to recombine enriched encoding templates.

In another embodiment of the invention when binding molecules against specific antigens that is only possible to express on a cell surface, e.g. ion channels or transmembrane receptors, is required, the cells particle themselves can be used as the selection agent. In this sort of approach, cells lacking the specific target should be used to do one or more rounds of negative selection or be present in large excass in the selection process. Here, irrelevant template-displayed molecules are removed. For example, for a positive selection against a receptor expressed on whole cells, the negative selection would be against the untransformed cells. This approach is also called subtraction selection and has successfully been used for phage display on antibody libraries (Hoogenboom et al. (1998) Immunotech. 4: 1-

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A specific example of a selection procedure can involve selection against cell surface receptors that become internalized from the membrane so that the receptor together with the selected binding molecule can make its way into the cell cytoplasm or cell nucleus. Depending on the dissociation rate constant for specific selected binding molecules, these molecules largely reside after uptake in either the cytoplasm or the nucleus.

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The skilled person in the art will acknowledge that the selection process can be per20 formed in any setup where the target is used as the bait onto which the templatedisplaying molecules can coordinate.

The selection methods of the present invention can be combined with secondary selection or screening to identify molecule ligands capable of modifying target molecule function upon binding. Thus, the methods described herein can be employed to isolate or produce binding molecules which bind to and modify the function of any protein or nucleic acid. It is contemplated that the method of the present invention can be employed to identify, isolate or produce binding molecules which will affect catalytic activity of target enzymes, i.e., inhibit catalysis or modifying substrate binding, affect the functionality of protein receptors, i.e., inhibit binding to receptors or modify the specificity of binding to receptors; affect the formation of protein multimers, i.e., disrupt quatemary structure of protein subunits; and modify transport properties of protein, i.e., disrupt transport of small molecules or ions by proteins.

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A still further aspect of the present invention relates to methods allowing functionality in the selection process can also be included. For example, when emichment against a certain target have been performed generation a number of different hits, these hits can then directly be tested for functionality (e.g. cell signalling). This can

for example be performed using fluorescence-activated cell sorting (FACS).

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The altered phenotype may be detected in a wide variety of ways. Generally, the changed phenotype is detected using, for example: microscopic analysis of cell morphology; standard cell viability assays, including both increased cell death and increased cell viability; standard labelling assays such as fluorometric indicator assays for the presence of level of particular cell or molecule, including FACS or other dye staining techniques; biochemical detection of the expression of target compounds after killing the cells; etc. In some cases, specific signalling pathways can be probed using various reporter gene constructs.

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Secondary selection methods that can be combined with template-displaying molecule technology include among others selections or screens for enzyme inhibition, alteration or substrate binding, loss of functionality, disruption of structure, etc.

Those of ordinary skill in the art are able to select among various alternatives of se-

ection or screening methods that are compatible with the methods described herein.

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The binding molecules of the invention can be selected for other properties in addition to binding. For example, during selection; stability to certain conditions of the desired working environment of the end product can be included as a selection crite-

desired, that protease can be part of the buffer medium used during selection. Similarly, the selection can also be performed in serum or cell extracts or any type of media. As will be understood, when utilizing this template-displaying approach, conditions which disrupt or degrade the template should be avoided to allow amplification. Other desired properties can be incorporated, directly into the displaying molecules as will be understood by those skilled in the art. For example, membrane affinity can be included as a property by employing building blocks with high hydropho-

duced by various synthetic methods. Chemical synthesis can be accomplished since the structure of selected binding molecules is readily obtained form the nucleic acid sequence of the coding template. Chemical synthesis of the selected molecules is also possible because the building blocks that compose the binding molecules are Molecules selected by the template-displaying molecule technology can be proaiso known in addition to the chemical reactions that assemble them together.

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tested in various appropriate in vitro and in vivo testing to verify the selected candidates for biological effects and potency. This may be done in a variety of ways, as will be appreciated by those in the art, and may depend on the composition of the In a preferred embodiment, the selected binding molecules is synthesized and bioactive molecule.

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Target identification and validation

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molecule technology could be used to select for specific epitopes on antigens found aspect, the target molecules are again preferably proteins or nucleic acids, but can also include, among others, carbohydrates and various molecules to which specific on cells, tissues or in vivo. These epitopes might belong to a target that is involved in important biological events. In addition, these epitopes might also be involved in In another aspect, the present invention provides methods to Identify or isolate targets that are involved in pathological processes or other biological events. In this molecule ligand binding can be achieved. In principal, the template-displaying the biological function of the target.

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successfully in Identifying new cellular antigens. (e.g. Pasqualini et al. (1996) Nature lecting for cell-surface antigen, the template molecule can be maintained outside the Phage display with antibodies and peptide libraries has been used numerous times (2002) Br J Cancer 86: 954-962; Kupsch et al. (1999) Clin Cancer Res. 5: 925-931; directly on cells suspected to express cell-specific antigens. Importantly, when secell. This will increase the probability that the template molecule will be intact after rseng-Law et al. (1999) Exp. Hematol. 27: 936-945; Gevorkian et al. (1998) Clin. Immunol. Immunopathol. 86: 305-309). Especially effective have been selection 380: 364-366; Pasqualini et al. (2000) Cancer Res. 60: 722-727; Scheffer et al. release for the cell surface.

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lecting from libraries of template-displayed molecules in vivo it is possible to isolate n vivo selection of temptate-displayed molecules has tremendous potential. By semolecules capable of homing specifically to normal tissues and other pathological issues (e.g. tumours). This principle has been illustrated using phage display of

- organs (Arap et al. (2002) Nat. Med. 2:121-127). A similar selection procedure could protected effectively by the phage particle allows selection in vivo. Accordingly, the peptide libraries (Pasqualini & Ruoslathi (1996) Nature 280: 364-366). This system be used for the template-displaying libraries. The coding DNA in phage display is has also been used in humans to identify peptide motifs that localized to different S
- ture Biotechnol. 14: 1116-1121; Pagratis et al. (1997) Nature Biotechnol. 15: 68-72). The template can be stabilised using various nucleotide derivatives in a similar way stability of the template in vivo will be important for amplification and identification. as have been used to stabilise aptamers for in vivo applications (Nolte (1996) Na-However, it is reasonable to believe that the template structure will be stabilized \$ 2
 - molecule. Other types of protection are also possible where the template molecule is shielded for the solution using various methods. This could include for example lipomolecules on the outside. The arrangement will protect the template molecules from plate form external manipulation. Fort example, the linker can be design to be incorsomes, pegylation, binding proteins or other sorts of protection. The template molecule could also be integrated into another designed structure that protects the temporated in vesicles to position the templates inside the vesicle and the displaying external manipulate and at the same time allow exposure of the displaying moleagainst degradation due to the modified bases used for encoding the displayed

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antigens (e.g. on a cell surface). The template-displaying technology should be able molecule (150 KDa) which will sterically reduce the access for a number of different Most antibodies have a large concave binding area which requires to some degree protruding epitopes on the antigens. Also, the antibody molecule is a large macro-

sules to permit selection.

- molecules will be able to bind into active sites, grooves and other areas on an antidiversity and complexity of the template-displaying molecule libraries will be much crease the physical access of the template-binding molecule par. In addition, the gen. The coding template element is also smaller that an antibody which will into access and recognize epitopes inaccessible to antibodies. The small binding ဓ
- greater compare to peptide libraries. This will increase the possibility to find mole-32

ides. One of ordinary skill in the art will acknowledge that various types of cells can new antigens can be performed using subtraction methods as described previously. tial to identify novel antigens which is not possible to identify with antibodies or pepchemistry. All together, the template-displaying molecule technology has the potenbe used in the selection procedure. It will also be understood that the selection for cules that can coordinate to epitopes inaccessible to peptides due to Inadequate

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cell-based assay or directly in vivo studying any phenotypic response. The strength dation of various targets. Most favourable, the binding molecules could also directly of this approach is that the same motecules are used both for identification and vali-Another aspect of the present invention relates to methods to validate the identified target. The identified binding molecules can directly be used if they change the blological response of the target. This can be done either in vitro using any direct or be used as therapeutic agents.

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In another preferred embodiment, the template-displaying molecules are used to pull plate-displaying molecule library with a cDNA library it will be possible to find binding The selected library is then plated to localized phage clones and the DNA coding for proteins from the cDNA library. One possibility is to mix a phage display library with a template display library and do a selection for either the phage or template library. Other types of libraries than cDNA could also be used such as nucleic aclds, carbopairs between the small molecules in the template-displaying molecule library and cDNA library expressed on bacteriophage (libraries vs. libraries). By mixing a temout the target molecules. This can for instance be achieved by selection against a the phage and template displayed molecules can then be identified using PCR.

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both phase I (activation) and phase II (detoxification) reactions. The major classes of corresponding to these displayed molecules could subsequently be used to compete in another embodiment of the invention the template-displaying molecule technology reactions are oxidation, reduction, and hydrolysls. Other enzymes catalyze conjugacan be used to account for in vivo and in vitro drug metabolism. That could include tions. These enzymes could be used as targets in a selection process to eliminate displayed molecule that are prone to coordinate to these enzymes. The templates or eliminate these molecules when making template-displaying molecule libraries. nydrates, synthetic polymer.

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binding to enzymes involved in phase I-II and possible be faster eliminated. For in-P450 enzymes, flavin monooxygenase, monoamine oxidase, esterases, amidases, These obtained libraries will then be free of molecules that will have a tendency of stance, a selection on each separate enzyme or any combination of cytochrome

glutathione S-transferases as well as other relevant enzymes could be performed to nydrolases, reductases, dehydrogenases, oxidases UDP-glucuronosyltransferases, identify these binding molecules that are prone to coordinate to these metabolic ensymes. Inhibitors are easily selected for due to their binding affinity but substrates need at least micro molar affinity to be identified. 2

good model for the epithelial barrier in the gastrointestinal guts. The CaCO-2 assay Another interesting embodiment of this invention is the possibility to directly select CaCO-2 cells, a human colon epithelial cell line, which is general, accepted as a plasma membrane, or other membranes. One possible selection assay is to use or molecules that passively or actively becomes transported across epithelial 5

solateral compartments. The template-displaying molecule libraries are placed either is collected and amplified. This process can be repeated until active molecules have such that the resultant monolayer forms a biological barrier between apical and baside of the cell monolayer and the molecules that can permeate the cell monolayer been identified. Other cell line or setup of this assay is possible and is obvious for nvolves growing a human colon epithelial cell line on tissue culture well inserts,

skill in the art.

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A still further aspect of the present invention relates methods of selecting for stability structure of the binding molecules. Various conditions could be certain proteases or a mixture of protease, cell extract, and various fluids from for example the gastroinof the selected molecules. This could be performed by subjecting an enriched pool that library to stability tests and selection to identify stable molecules under certain testinal gut. Other conditions could be various salts or acld milieu or elevated temperature. Another possibility is to generate a library of known ligands and subject of binding molecules to an environment that will possibly degrade or change the

Therapeutic applications

conditions as describe above.

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known or suspected to be involved in a regulating process that is malfunctioning and or stimulating various targets. A therapeutically relevant target is a substance that is template-displaying molecule technology of the invention may be used for blocking molecules, cofactor-enzyme interaction, and protein-protein interaction in intracellucule ligand is desired. Thus, target can, for example, include a chemical compound, library, a ribosome peptide display library, an extract made from blological materials a mixture of chemical compounds, an array of spatially localized compounds, a bloteraction, transcription-DNA interaction, and cell-cell interaction involving adhesion fusion protein, peptide, enzyme, receptor, receptor ligand, hormone, antigen, anti-The potential therapeutic applications of the invention are great. For example, the thus leads to a disease state. Examples of such processes are receptor-ligand inlar signalling. Target molecule means any compound of interest for which a molesuch as bacteria, plants, fungi, or animal (e.g. mammalian) cells or tissue, protein, body, drug, dye, growth factor, lipid, substrate, toxin, virus, or the like etc., without mixture of related or unrelated proteins, a mixture of viruses or bacterial strains or limitation. Other examples of targets include, e.g. a whole cell, a whole tissue, a logical macromolecule, such as DNA or mRNA, a bacteriophage peptide display the like. etc., without limitation.

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ases for pharmacological intervention. Although the above examples are focused on nuclear receptors, and metabolic enzymes constitute overwhelmingly the majority of the most relevant targets, it will be self-evident for a person skilled in the art that any Therapeutic drug targets can be divided into different classes according to function; known targets for existing drugs. Especially, G Protein-Coupled Receptors (GPCR) receptors, enzymes, hormones, transcription factors, ion channels, nuclear recepconstitutes one of the most important classes of drug targets together with protetors, DNA, (Drews, J. (2000) Science 287:1960-1964). Among those, receptors, other therapeutic target may be of Interest.

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dependent on the specific properties each target holds. Most of the targets are possible to obtain in a purified form for direct selection procedures. Other targets have The present invention employing the template-displaying molecule technology can to be used when they are in their native environments such as imbedded cell surbe utilized to identify agonists or antagonists for all these classes of drug targets,

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face receptors. In those situations the selection using the template-displaying molecule libraries can be performed using subtraction-selection described previously.

volving antiviral agents are also included. For example, a generated molecule, which Such molecules may in addition carry another therapeutic agent to increase the poapplication includes cell targeting. For example, the generated molecules recognizbinds strongly to epitopes on the virus particle, may be useful as an antiviral agent. tency and reduce the side-effects (for example cancer treatment). Applications in-Another specific application of the template-displaying molecule technology of the vention is to generate molecules that can function as antagonists, where the mole-One specific application of the template-displaying molecule technology of the ining specific surface proteins or receptors will be able to bind to certain cell types. invention is to generate molecules that can function as agonists, where the molecules block the interaction between a receptor and one or more ligands. Another cules stimulate or activate a receptor to initiate a cellular signalling pathway. S 2 5

femplate-displaying molecule arrays

lescribed herein. These molecule ligands can be used separately or in array system sample, which employs a molecule ligand which can be isolated by the methods A still further aspect of the present invention relates to methods for detecting the presence or absence of, and for measuring the amount of target molecules in a for multiple determinations.

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An understanding of protein structures, protein-to-protein interactions, pathways and rays. The protein microarrays would be very sensitive to and can be easily degraded much more difficult. Unlike hybridization reactions, which are based on couplings or how proteins influence the origins of disease is of vital importance. Nucleic acid ming. However, a major hurdle is the lack of correlation between gene expression at the level of mRNA level and the amount of corresponding protein expressed within the cell (Andersson et a. (1997) Electrophiresis 18: 533-537). Contrary to DNA and croarrays have enabled researchers to pursue novel biomarkers through genotypaces anising from 3D folded amino-acid sequences. The requirement for preparation of 3D folded protelns substantially complicates fabrication of protein microarinteractions of linear sequences, the protein interactions involve polypeptide sur-RNA analysis, the use of biochips for parallel protein function studies has been

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bridization reactions used on the DNA/RNA biochlps. The sequence dependency of by the use of thermal treatments and harsh chemicals. Moreover, the folded protein interactions have a much stronger dependency on sequences compared to the hythe protein interactions will further complicate the reaction kinetics.

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cule against cytokines or enzymes known to be involved in a specific pathway could molecule to measure the amount of various biomarkers. For example, binding molespotted in an array format to be used to measure the absolute or relative amount of could be used to identify small binding molecules to numerous targets. These bindbe generate with the describe technology. These binding molecules could then be The invention described herein provides a possible solution to making arrays that can measure different amounts at the protein level without the use of proteins or ing molecules could then be arrayed in specific positions and work as detection peptides as detection molecules. The template-displaying molecule technology each cytokine or enzyme.

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One major advantage with this system is that the spotting technology used for DNA tional molecules onto different locations of a chip. The overall principal is shown in arrays could be identically applied for this system. The template-displayed moletechnology will lead to high-throughput deposition of thousands of different funccules could be directly applied to the spotted DNA. Another possibility is that the merase and the nucleotide analogues. Make addressable microarrays with this synthesis could be performed directly on the pre-coated template using a polyfigure 40.

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molecules will be more suitable to become immobilized on a surface and exposed to any harsh conditions such as heat, tow or high pH various detergent. In addition, the The template-displaying molecule technology is not limited in chemistry to the 20 natural occurring amino acids. This will permit synthesis on the template of more robust and stable molecules that will bind to various targets. These more stable shelf-life of the arrays will be much longer that arrays made from proteins.

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Molecular biological tools

Polymerase chain reaction (PCR) is an exemplary method for amplifying nucleic acids. Descriptions of PCR methods are found (Saiki et al. (1985) Science 230:

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(Mullis et al.)). Alternative methods of amplification include among others cloning of that will allow faithful, efficient amplification of selected nucleic acid sequences can (Guatelli et al. (1990) Proc. Natl. Acad, Scl. 87: 1874-1878). In general, any means organism where the vector and the cloned DNAs are replicated and thus amplified be employed in the method of the present invention. It is only necessary that the 1350-1354; Scharf et al. (1986) Science 233; 1076-1078; U.S Patent 4,683,202 selected DNAs into appropriate vector and introduction of that vector into a host

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proportionate representations of the sequences after amplification reflect the relative

proportions of sequences in the mixture before amplification.

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by chemical synthests or a combination of chemical synthests and recombinant DNA method known in the art. Such methods include constructing a nucleotide sequence The template variants of the present invention may be produced by any suitable technology.

introduction (I.e. Insertion or substitution) or removal (I.e. deletion or substitution) of structed by isolating or synthesizing a nucleotide sequence encoding the appropriate display molecules and then changing the nucleotide sequence so as to effect A nucleotide sequence encoding a template variant of the invention may be conthe relevant functional entities of the displayed molecutes.

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ing for portions of the desIred template may be synthesized and assembled by PCR, igation or ligation chain reaction (LCR) (Barany, PNAS 88:189-193, 1991). The indion the sequence of the desired templates. For example, small oligonucleotides codusing an oligonucleotide synthesizer, wherein oligonucleotides are designed based The nucleotide sequence corresponding to the template molecules is conveniently Alternatively, the nucleotide sequence is prepared by chemical synthesis, e.g. by vidual oligonucleotides typically contain 5' or 3' overhangs for complementary asmodified by site-directed mutagenesis in accordance with conventional methods.

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template variants for high throughput screening or selection. For instance, methods Alternative nucleotide sequence modification methods are available for producing which involve homologous cross-over such as disclosed in US 5,093,257, and

methods which involve gene shuffling, i.e. recombination between two or more ho-32

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mologous nucleotide sequences resulting in new nucleotide sequences having a number of nucleotide alterations when compared to the starting nucleotide sequences. Gene shuffling (also known as DNA shuffling) involves one or more cycles of random fragmentation and reassembly of the nucleotide sequences, followed by selection to select nucleotide template sequences encoding variant displaying molecules with the desired properties.

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Examples of sultable in vitro gene shuffiling methods are disclosed by Stemmer et al. (1994), Proc. Natl. Acad. Sci. USA; vol. 91, pp. 10747-10751; Stemmer (1994), Nature, vol. 370, pp. 389-391; Smith (1994), Nature vol. 370, pp. 324-325; Zhao et al., Nat. Biotechnol. 1998, Mar. 16(3); 258-61; Zhao H. and Amold, FB, Nucleic Acids Research, 1997, Vol. 25. No. 6 pp. 1307-1308; Shao et al., Nucleic Acids Research 1998, Jan 15; 26(2); pp. 681-83; and WO 95/17413.

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Synthetic shuffling involves providing libraries of overlapping synthetic oligonucleotides based e.g. on a flanking sequence. The synthetically generated oligonucleotides are recombined, and the resulting recombinant nucleic acid sequences are screened and if desired used for further shuffling cycles.

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Recombination can be theoretically calculated, which is performed or modelled using a computer system, thereby partly or entirely avoiding the need for physically manipulating nucleic acids.

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Once assembled (by synthesis, site-directed mutagenesis, DNA shuffiling or another method), the nucleotide sequence encoding the templates is used to generate the template-displaying libraries.

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Still other aspects of the present invention relates to a pharmaceutical composition comprising the conjugate or the variant of the invention as well as to methods of producing and using the conjugates and variants of the invention.

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The term "affinity" is used herein as a qualitative term to describe the molecule-target Interaction. A quantitative measure for the affinity is expressed through the Association Constant and the Dissociation Constant is related to each other by the equation K_o = 1/K_o. Evidently, a high affinity corre-

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sponds to a lower Dissociation Constant. The term "binds to a specific targer" means that the binding molecules obtained with the template-displaying molecule technology binds to a chosen target so that a measurable response is obtained when tested in a suitable binding or functional assay. In the present context, the term "therapeutic agent" is intended to mean any biologically or pharmacologically active substance or antigen-comprising material; the term includes substances which have utility in the treatment or prevention of diseases or disorders affecting animals and humans, or in the regulation of airy animal or human physiological condition and it also includes any biological active compound or composition which, when administrated in an effective amount, has an effect on living cells or organisms.

and incubate for about 30 min at 20 °C and then remove the excess by washing with wells of two flat-bottom microtiter plates with about 1 µg streptavIdIn in a TBS buffer. After the construction of template-displayed libraries, template-displaying molecules incubate over night at 4 °C. Remove the streptavidin solution and wash the wells at or about 30 min. at 37 °C. Wash the plate with TBS buffer at least three times. Add plate-displaying molecule library to both wells and allow binding by incubating at 20 BS buffer at least six times. Block free streptavidin molecules with 1 mM biotin for 5 min. and wash excess away with TBS buffer at least six times. Add then the temleast six times with TBS. Immediately add 2% BSA to block the wells and incubate cule. Elute the coordinated template-displaying molecules using condition that rescribed in the literature) to one of the wells (use the other as background control) emplate-displaying molecules that not coordinate to the immobilized target mole-°C for about 1 hour. Wash the wells with TBS buffer at least six times to remove bearing the desired ligands can be captured using the below protocol. Coat two about 0.1 µg biotinylated target molecule (biotinylation can be performed as de-र्ठ ឧ 2

molecules between the wells with and without the target molecule to make sure there are more template-displaying molecules eluted in the well with target. That will ensure that there is a specific enrichment in the selection process. Other types and numerous variations of selection procedures can be found in the literature (e.g. "Phage display: A laboratory manual" (2001) Barbas et al., Eds. Cold Spring Harbor Laboratory Press, New York),

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move the binding molecules. In later selection cycles, compare the number of eluted

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Methods 206: 73-85). Positive cells (cells with the antigen of interest) is cell-surface An alternative to the above capturing is, after the construction of template-displayed ing. Add about 10⁸ negative cells (cells without the antigen of interest). These negacells to 10 µl streptavidin-coated paramagnetic microbeads (Dynal) and allow bindcells capture the specific template-displaying molecules. Pellet the cell mixture, disbiotinylated using sulfo-NHS-LC-biotin (Pierce). Add approximately 108 biotinylated positive cells by wash off all the negative cells. Finally elute the positive cells by removing the magnetic field and amplify the eluted templates using PCR. This selective cells act as a sink for nonpecific template-displaying molecules, and the target Incubate about 2 hours at 37 °C on a rotator to keep the cells in suspension. Load libraries, to capture the template-displaying molecules bearing the desired ligands using the below protocol. The selection of template-display molecules can be percard the supernatant, and suspend in the template-displaying library suspension. the cell/template-displaying library solution on a magnetic column to recover the formed using magnetically activated cell sorting (Siegel et al. (1997) J. Immunol. lion protocol can be repeated several times if needed.

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Amplification of templates capable of templating the synthesis of templated molecules

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In one aspect the present invention relates to methods for amplifying templated molecules that may or may not be bound to a target. The choice of amplification method depends on the choice of coding or complementing elements. Natural oligonucleotides can be amplified by any state of the art method. These methods include, but is not limited to the polymerase chain reaction (PCR); as wells as e.g. nucleic acid sequence-based amplification (e.g. Compton, Nature 350, 91-92 (1991)), amplified anti-sense RNA (e.g. van Gelder et al., PNAS 85: 77652-77666 (1988)); self-sustained sequence replication system (e.g. Gnatelli et al., PNAS 87: 1874-1878 (1990)); polymerase independent amplification as described in e.g. Schmidt et al., NAR 25: 4797-4802 (1997), as well as in vivo amplification of plasmids carrying cloned DNA fragments. Ligase-mediated amplification methods may also be used, e.g., LCR (Ligase Chain Reaction).

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For non-natural nucleotides the choices of efficient amplification procedures are fewer. As non-natural nucleotides per definition can be incorporated by certain enzymes including polymerases, it will be possible to perform manual polymerase chain reaction by adding the polymerase during each extension cycle.

For oligonucleotides containing nucleotide analogs, fewer methods for amplification

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exist. One may use non-enzyme mediated amplification schemes (Schmidt et al., NAR 25: 4797-4802 (1997)). For backbone-modified oligonucleotide analogs such as PNA and LNA, this amplification method may be used. Before or during amplification the templates or complementing templates may be mutagenized or recombined in order to create a larger diversity for the next round of selection or

Characterization of polymers isolated by the selections or screening assays.

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After the final round of selection, it is often desirable to sequence individual templates, in order to determine the sequence of individual templated polymers. If the template contains natural nucleotides, it is a standard routine to optionally PCR amplify the isolated templates (if the template is an RNA molecule, it is necessary to use reverse transcriptase to produce cDNA prior to the PCR-amplification), and then clone the DNA fragments into for example plasmids, transform these and then sequence individual plasmid-clones containing one or multiple tandem DNA sequences. In this case, it is practical to design a restriction site in both of the flanking sequences to the central random or partly random sequence of the template (i.e., in the primer binding sites). This will allow easy cloning of the isolated nucleotides. Sequencing can be done by the standard dideoxy chain termination method, or by

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If the template contains non-natural nucleotides, it is not feasible to clone individual sequences by transfer through a microbial host. However, using bead populations where each bead carries one oligonucleotide sequence, it is possible to clone in vitro, whereafter all the nucleotides ettached to a specific bead may be optionally amplified and then sequenced (Brenner et al., 2000, Proc. Natl. Acad. Sci. USA 97, 1665-1670). Alternatively, one may dilute the population of isolates adequately, and

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nore classical means such as Maxam-Gilbert sequencing.

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non-natural nucleotides are substrates for the thermostable polymerase used in the be possible to sequence using standard methods. Of course, this requires that the then aliquot into microtiter plates so that the wells on average contain for example 0.1 templates. By amplifying the single templates by for example PCR, it will now

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able to design the starting template so as to contain restriction sites on either side of If alternative methods are used that require shorter ofigonucleotides it may be desirthe encoding/templating region of the template. Thereby, after the final selection

round, the templates can be restricted, to obtain a short oligonucleotide encoding the templated polymer, and then these short oligos can be applied to various anaytical procedures. 5

It is also possible to sequence the isolates by the use of a DNA array of oligos with 5

random but predetermined sequences.

It may also be desirable to sequence the population of isolates as a pool, for example if the sequences are expected to be in register, for example because the initial lection. Wherefore the population of isolates can be sequenced as a whole, to obsolates have sequences similar to the initial sequence of the templates before selibrary consisted of a degenerate sequence based on a polymer sequence with a known (relatively high) desired activity. Therefore, it is then expected that all the tain a consensus sequence for the population as a whole.

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Templated molecules 23

A non-exhaustive and non-limiting list of oligomers that may be templated by the various principles described in the present invention is listed below:

alpha-, beta-, gamma-, and omega-peptides

mono-, di- and tri-substituted peptides

cyclohexane- and cyclopentane-backbone modified beta-peptides L- and D-form peptides

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vinylogous polypeptides glycopolypeptides

polyamides

vinylogous sulfonamide peptide

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conjugated peptide (i.e., having prosthetic groups) Polysulfonamide

Polyesters

Polysaccharides

Polycarbamates

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Polycarbonates

Polyureas

poly-peptidy/phosphonates

Azatides

Polyethers

polyethylene glycols (PEG)

Polyethylenes

Polydisulfides

oolyarylene suifides

Polynucleotides

PNAs

Morpholinos

Polyethylenelmine

22

Polyacetates

Polystyrenes

8

Glycolipids

polycycles (aromatic)

polyheterocycles 33

peptoids (oligo N-substituted glycines) 9

ethoxyformacetal oligomers

poly-thioethers

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LNAs

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oligo pyrrolinone polyoximes

Polyimines

olyacetylene

Polyvinyl

Phospholipids Lipids

polycycles (aliphatic)

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Polysiloxanes Proteoglycan

Polyisocyanides

Polyisocyanates

Polymethacryfates

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Monofunctional, Difunctional, Trifunctional and Oligofunctional open-chain hydrocarbons.

Monofunctional, Difunctional, Trifunctional and Oligofunctional Nonaromatic

Carbocycles.

Monocyclic, Bicyclic, Tricyclic and Polycyclic Hydrocarbons 2

Bridged Polycyclic Hydrocarbones

Monofunctional, Difunctional, Trifunctional and Oligofunctional Nonaromatic

Heterocycles.

Monocyclic, Bicyclic, Tricyclic and Polycyclic Heterocycles

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Bridged Polycyclic Heterocycles

Monofunctional, Difunctional, Trifunctional and Oligofunctional Aromatic Carbo-

cycles.

Monocyclic, Bicyclic, Tricyclic and Polycyclic Aromatic Carbocycles

Monofunctional, Difunctional, Trifunctional and Oligofunctional Aromatic Hetero-

cycles. 8 Monocyclic, Bicyclic, Tricyclic and Polycyclic Heterocycles

Chelates

Fullerenes.

Any combination of the above.

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The list refers to any linear, branched or cyclic structure that contains one or more of the backbone structures listed, and/or contain several bonds of the same kind (e.g. amide bonds). Heteropolymers (hybrids of different polymer types) can also be lemplated by the present invention.

Below a table is presented stating the polymers producible according to the present invention as well as the functional entities/reactive groups required to make them. A reference is made to the relevant figure:

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Fig. 11, 6x. 1 in 15 Fin 15, ex. E E Catalysureagent Sahne conditions Ilkylating agent glycogen synihetese polycaccharide syniheteses base EDC or other carbodimide ethodimide carbodimide inking malacula casbonyldilmidazole ativation system (Kohne stycosylation) mine, N-ydroxysuscinimide ester cahal, carboxylic seld alonen, carboxyfio acid uning, carboxylic acid Icobel, Isocyanata condary amina, o uppropriest forthe Functional Entity (reactive groups) ysaccharide olyanschaufte lycarbonate ohacetale olyacetala oppuesto polyamida

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aplyagailde Softwaride. pohypeptida aplicantide molyne pittle

actrophida

polyamide

Phosphonale

potrohosotate.

Yohosphonele

Fig. 22, Fig. 22, Fig. 15, ex. 1 Fig. 15, 84, 7 Ffn. 18 Pfg. 18, ex. 1 Fig. 18, 0x.2 Fq. 11 Fa. 15 Fib. 18 General Linking mulecula Catalyst/reagant Figura Giscorgante Fig. 19 Fo. 44 운 다 다 다 유럽 건경 Fb, 13 radioal initiator, AIBN di-sikone (henzoquinone) mckostás Siphosphorooldehyde, hydronylamine iming, alkyl sulfangte aldehyde, emine aldehyde, emine Functional Endby (rozedvo groups) diamina shylchloride unit thlol, thiol hicepoxida styrene-und di-Chane akene polynacleatides polycycloalkane pohydasufide poptraoether potyethylene polystyrene polyovime polymine polymine polyscrine

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Templates

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molecule that is not linked to the complementing template or template that templated templated molecule. In another embodiment, the method for templating a templated In one embodiment, the templated molecule is linked by means of a single linker to molecule comprises the further step of releasing the template or complementing the complementing template or template that templated the synthesis of the template that templated the templated molecule, and obtaining a templated the synthesis of the templated molecule.

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such as from 3 to 20, such as from 3 to 15, for example from 3 to 15, such as from 3 for example from 4 to 10, such as from 4 to 8, such as from 4 to 6, for example 4, for example from 5 to 100, such as from 5 to 80, for example from 5 to 60, such as from to 10, such as from 3 to 8, for example from 3 to 6, such as from 3 to 4, for example such as from 6 to 100, for example from 6 to 80, such as from 6 to 60, such as from such as from 7 to 8, for example 7, for example from 8 to 100, such as from 8 to 80, from 8 to 20, for example from 8 to 15, such as from 8 to 10, such as 8, for example value of from 2 to 200, for example from 2 to 100, such as from 2 to 80, for example from 2 to 6, such as from 2 to 4, for example 2, such as from 3 to 100, for example from 4 to 40, for example from 4 to 30, such as from 4 to 20, such as from 4 to 15, from 7 to 30, such as from 7 to 20, for example from 7 to 15, such as from 7 to 10, from 2 to 60, such as from 2 to 40, for example from 2 to 30, such as from 2 to 20, from 3 to 80, such as from 3 to 60, such as from 3 to 40, for example from 3 to 30, 3, such as from 4 to 100, for example from 4 to 80, such as from 4 to 60, such as such as from 5 to 10, such as from 5 to 8, for example from 5 to 6, for example 5, such as from 7 to 80, for example from 7 to 60, such as from 7 to 40, for example for example from 8 to 60, such as from 8 to 40, for example from 8 to 30, such as template comprising n coding elements can also be branched. n preferably has a 5 to 40, for example from 5 to 30, such as from 5 to 20, for example from 5 to 15, 6 to 40, for example from 6 to 30, such as from 6 to 20, such as from 6 to 15, for example from 6 to 10, such as from 6 to 8, such as 6, for example from 7 to 100, for example from 2 to 15, such as from 2 to 10, such as from 2 to 8, for example 9, for example from 10 to 100, such as from 10 to 80, for example from 10 to 60, The template preferably comprises n coding elements in a linear sequence. The

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example from 12 to 30, such as from 12 to 20, for example from 12 to 15, such as example from 10 to 15, such as from 10 to 12, such as 10, for example from 12 to 100, such as from 12 to 80, for example from 12 to 60, such as from 12 to 40, for such as from 10 to 40, for example from 10 to 30, such as from 10 to 20, for

- from 14 to 100, such as from 14 to 80, for example from 14 to 60, such as from 14 to from 16 to 40, for example from 16 to 30, such as from 16 to 20, such as from 18 to such as from 16 to 100, such as from 16 to 80, for example from 16 to 60, such as 100, such as from 18 to 80, for example from 18 to 60, such as from 18 to 40, for 10, for example from 14 to 30, such as from 14 to 20, for example from 14 to 16, z,
- example from 18 to 30, such as from 18 to 20, for example from 20 to 100, such as 20 to 30, such as from 20 to 25, for example from 22 to 100, such as from 22 to 80, for example from 22 to 60, such as from 22 to 40, for example from 22 to 30, such from 20 to 80, for example from 20 to 60, such as from 20 to 40, for example from as from 22 to 25, for example from 25 to 100, such as from 25 to 80, for example 9
- from 25 to 60, such as from 25 to 40, for example from 25 to 30, such as from 30 to example from 35 to 60, such as from 35 to 40, for example from 40 to 100, such as from 40 to 80, for example from 40 to 60, such as from 40 to 50, for example from 100, for example from 30 to 80, such as from 30 to 60, for example from 30 to 40, such as from 30 to 35, for example from 35 to 100, such as from 35 to 80, for 15
- from 60 to 80, such as from 60 to 70, for example from 70 to 100, such as from 70 to for example from 45 to 50, such as from 50 to 100, for example from 50 to 80, such 40 to 45, such as from 45 to 100, for example from 45 to 80, such as from 45 to 60, 90, for example from 70 to 80, such as from 80 to 100, for example from 80 to 90, as from 50 to 60, for example from 50 to 55, such as from 60 to 100, for example ន
- such as from 90 to 100. 22

In some embodiments of the invention it is preferred that the template is attached to a solid or semi-solid support.

ribonucleic acids (RNA), peptide nucleic acids (PNA), locked nucleic acids (LNA), The template in one embodiment preferably comprises or essentially consists of nucleotides selected from the group consisting of deoxyribonucleic acids (DNA), and morpholinos sequences, including any analog or derivative thereof. ജ

In another embodiment, the template of coding elements preferably comprises or essentially consists of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof, and the complementing element preferably comprises or essentially consists of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof.

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It is preferred in various embodiments of the Invention that the template can be characterised by any one or more of the following features: I) That the template is amplifyable, ii) that the template comprises a single strand of coding elements, preferably a single strand of coding elements capable of forming a double helix by hybridization to a complementing template comprising a single strand of complementing template comprises a priming site.

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15 Coding elements

Each coding element is preferably linked to a neighbouring coding element by a covalent chemical bond. Each coding element can also be linked to each neighbouring coding element by a covalent chemical bond. The covalent chemical bond is preferably selected from the group of covalent bonds consisting of phosphodiester bonds, phosphorothioate bonds, and peptide bonds. More preferably, the covalent chemical bond is selected from the group of covalent bonds consisting of phosphodiester bonds and phosphorothioate bonds.

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25 In preferred embodiments, at least one coding element is attached to a solid or semi-solid support. The coding elements are selected in one embodiment of the inveniton from the

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group consisting of nucleotides, including any analog or derivative thereof, amino acids, antibodies, and antigens, and preferably from the group consisting of nucleotides, nucleotide derivatives, and nucleotide analogs, including any combination thereof. In another embodiment, the coding elements are selected from the group consisting of nucleotides, including nucleotides such as deoxyribonucleic acids comprising a base selected from adenine (A), thymine (T), guanine (G), and cytosine (C), and ribonucleic acids comprising a base selected from adenine (A),

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uracil (U), guanine (G), and cytosine (C). Also in this case can each nucleotide be linked to a neighbouring nucleotide by means of a covalent bond, or linked to each neighbouring nucleotide by means of a covalent bond. The covalent bond is preferably a phosphodiester bond or a phosphorothloate bond.

In other embodiments, the coding elements are natural and non-natural nucleotides selected from the group consisting of deoxyribonucleic acids and ribonucleic acids.

Coding Elements and Corresponding Complementing Elements

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When the coding elements are preferably selected from the group consisting of nucleotides, nucleotide derivatives and nucleotide analogs in which one or more of the base molety and/or the phosphate moiety and/or the ribose or deoxyribose molety have been substituted by an alternative molecular entity, corresponding complementing elements are capable of interacting with said coding elements and preferably comprise or essentially consist of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof. Each nucleotide is linked to a neighbouring nucleotide by a covalent chemical bond, or linked to each neighbouring nucleotide by a covalent chemical bond. The covalent chemical bond is preferably selected from the group of covalent bonds consisting of phosphodiester bonds and peptide bonds.

Coding Element Subunits

Coding elements in one embodiment preferably comprise or essentially consist of from 1 to 100 subunits, such as from 1 to 80 subunits, for example from 1 to 60 subunits, such as from 1 to 40 subunits, for example from 1 to 16 subunits, such as from 1 to 14 subunits, for example from 1 to 15 subunits, such as from 1 to 14 subunits, for example from 1 to 12 subunits, such as from 1 to 10 subunits, for example from 1 to 9 subunits, for example from 1 to 9 subunits, for example from 1 to 5 subunits, for example from 1 to 5 subunits, such as from 1 to 4 subunits, for example from 1 to 3 subunits, such as from 1 to 2 subunits, for example from 2 to 60 subunits, such as from 2 to 80 subunits, for example from 2 to 60 subunits, such as from 2 to 40 subunits, for example from 2 to 20 subunits, such as from 2 to 18 subunits, for example from 2 to 20 subunits, such as from 2 to 18 subunits, for example from 2 to 20 subunits, such as from 2 to 18 subunits, for example from 2 to 20 subunits, such as from 2 to 18 subunits, for example from 2 to 20 subunits, such as from 2 to 18 subunits, for example from 2 to 20 subunits, such as from 2 to 80 subunits, for example from 2 to 20 subunits, such as from 2 to 80 subunits, for example from 2 to 80 subunits, such 80 subunits, for example from 2 to 80 subunits, for example from 2 to 80 subunits, such 80 subunits, for example from 2 to 80 subunits, for example from 2 to 80 subunits, for example from 2 to 80 subunits, such 80 subunits for example from 2 to 80 subunits for 60 subunits

16 subunits, such as from 2 to 14 subunits, for example from 2 to 12 subunits, such

as from 2 to 10 subunits, for example from 2 to 9 subunits, such as from 2 to 8

subunits, for example from 2 to 7 subunits, such as from 2 to 6 subunits, for

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example from 8 to 20 subunits, such as from 8 to 18 subunits, for example from 8 to 16 subunits, such as from 8 to 14 subunits, for example from 8 to 12 subunits, such as from 8 to 10 subunits, for example from 8 to 9 subunits, for example 8 subunits, such as from 9 to 100 subunits, such as from 9 to 80 subunits, for example from 9 to

such as from 9 to 100 subunits, such as from 9 to 80 subunits, tor example from 9 to 60 subunits, such as from 9 to 40 subunits, for example from 9 to 20 subunits, such as from 9 to 18 subunits, for example from 9 to 16 subunits, such as from 9 to 14 subunits, for example from 9 to 12 subunits, such as from 9 to 10 subunits, such as 9 subunits, for example from 10 to 100 subunits, such as 6 subunits, for example from 10 to 60 subunits, such as from 10 to 40 subunits, for example from

example from 3 to 20 subunits, such as from 3 to 18 subunits, for example from 3 to

example from 2 to 5 subunits, such as from 2 to 4 subunits, for example from 2 to 3

subunits, such as 2 subunits, such as from 3 to 100 subunits, such as from 3 to 80

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subunits, for example from 3 to 60 subunits, such as from 3 to 40 subunits, for

16 subunits, such as from 3 to 14 subunits, for example from 3 to 12 subunits, such

as from 3 to 10 subunits, for example from 3 to 9 subunits, such as from 3 to 8

subunits, for example from 3 to 7 subunits, such as from 3 to 6 subunits, for

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for example from 4 to 100 subunits, such as from 4 to 80 subunits, for example from

example from 3 to 5 subunits, such as from 3 to 4 subunits, for example 3 subunits,

- 10 to 20 subunits, such as from 10 to 18 subunits, for example from 10 to 16 subunits, such as from 10 to 14 subunits, for example from 10 to 12 subunits, such as 10 subunits, such as from 11 to 100 subunits, such as from 11 to 60 subunits, such as from 11 to 40 subunits, for example from 11 to 20 subunits, such as from 11 to 18 subunits, for example from 11 to 20 subunits, such as from 11 to 18 subunits, for example from 11 to 20 subunits, such as from 11 to 18 subunits, for example from 11 to 16
- subunits, such as from 11 to 14 subunits, for example from 11 to 12 subunits, such as from 12 to 100 subunits, such as from 12 to 80 subunits, for example from 12 to 60 subunits, such as from 12 to 40 subunits, for example from 12 to 20 subunits, such as from 12 to 18 subunits, for example from 12 to 16 subunits, such as from 12 to 14 subunits, for example from 13 to 100 subunits, such as from 13 to 80 subunits,

for example from 5 to 60 subunits, such as from 5 to 40 subunits, for example from 5

such as from 5 to 14 subunits, for example from 5 to 12 subunits, such as from 5 to

10 subunits, for example from 5 to 9 subunits, such as from 5 to 8 subunits, for

to 20 subunits, such as from 5 to 18 subunits, for example from 5 to 16 subunits,

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example from 5 to 7 subunits, such as from 5 to 6 subunits, such as 5 subunits, for example from 6 to 100 subunits, such as from 6 to 80 subunits, for example from 6

example 4 subunits, such as from 5 to 100 subunits, such as from 5 to 80 subunits,

subunits, such as from 4 to 6 subunits, for example from 4 to 5 subunits, for

example from 4 to 9 subunits, such as from 4 to 8 subunits, for example from 4 to 7

such as from 4 to 18 subunits, for example from 4 to 16 subunits, such as from 4 to

14 subunits, for example from 4 to 12 subunits, such as from 4 to 10 subunits, for

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4 to 60 subunits, such as from 4 to 40 subunits, for example from 4 to 20 subunits,

- for example from 13 to 60 subunits, such as from 13 to 40 subunits, for example from 13 to 20 subunits, such as from 13 to 18 subunits, for example from 13 to 16 subunits, such as from 13 to 14 subunits, for example from 14 to 100 subunits, such as from 14 to 80 subunits, for example from 14 to 60 subunits, such as from 14 to 40 subunits, for example from 14 to 20 subunits, such as from 14 to 18 subunits, for
 - example from 14 to 16 subunits, such as from 15 to 100 subunits, such as from 15 to 80 subunits, for example from 15 to 60 subunits, such as from 15 to 40 subunits, for example from 15 to 20 subunits, such as from 15 to 18 subunits, for example from 15 to 16 subunits, such as from 16 to 100 subunits, such as from 16 to 80 subunits, for example from 16 to 20 subunits, such as from 16 to 40 subunits, for example from 17 to 100 subunits, such as from 16 to 18 subunits, for example from 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 60 subunits, such as from 17 to 80 subunits, for example from 17 to 60 subunits, such as from 17 to 80 subunits, for example from 17 to 60 subunits, such as from 17 to 80 subunits, for example from 17 to 60 subunits, such as from 17 to 80 subunits, such as from 17 to 80 subunits, such as from 17 to 60 subunits, such as from 17 to 80 subunits, such as from 17 to 60 subunits, such as from 17 to 80 subunits, such as from 17 to 60 subunits such as from 17 to 80 subunits such as from 17 to 60 subunits such as from 17 to 60 subunits such as from 17 to 80 subunits such as from 17 to 60 subunits su
- as from 17 to 18 subunits, for example from 18 to 100 subunits, such as from 18 to 80 subunits, for example from 18 to 60 subunits, such as from 18 to 40 subunits, for example from 18 to 20 subunits, such as from 19 to 100 subunits, such as from 19

subunits, such as from 17 to 40 subunits, for example from 17 to 20 subunits, such

example from 7 to 20 subunits, such as from 7 to 18 subunits, for example from 7 to

16 subunits, such as from 7 to 14 subunits, for example from 7 to 12 subunits, such

such as from 6 to 18 subunits, for example from 6 to 16 subunits, such as from 6 to

to 60 subunits, such as from 6 to 40 subunits, for example from 6 to 20 subunits,

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example from 6 to 9 subunits, such as from 6 to 8 subunits, for example from 6 to 7

14 subunits, for example from 6 to 12 subunits, such as from 6 to 10 subunits, for

subunits, such as 6 subunits, such as from 7 to 100 subunits, such as from 7 to 80

subunits, for example from 7 to 60 subunits, such as from 7 to 40 subunits, for

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subunits, such as 7 subunits, for example from 8 to 100 subunits, such as from 8 to

as from 7 to 10 subunits, for example from 7 to 9 subunits, such as from 7 to 8

80 subunits, for example from 8 to 60 subunits, such as from 8 to 40 subunits, for

from 20 to 100 subunits, such as from 20 to 80 subunits, for example from 20 to 60 to 80 subunits, for example from 19 to 60 subunits, such as from 19 to 40 subunits, subunits, such as from 20 to 40 subunits, for example from 20 to 30 subunits, such for example from 19 to 30 subunits, such as from 19 to 25 subunits, for example as from 20 to 25 subunits.

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a covalent bond, or linked to each neighbouring nucleotide, or nucleotide analog, by nucleotide is linked to a neighbouring nucleotide, or nucleotide analog, by means of consisting of phosphodiester bonds, phosphorothicate bonds, and peptide bonds. In preferred embodiments, each coding element subunit comprises or essentially deoxyribonucleic acid comprising a base selected from adenine (A), thymine (T), guanine (G), and cytosine (C), or it can be a ribonucleic acid comprising a base means of a covalent bond, including covalent bonds selected from the group selected from adenine (A), uracil (U), guanine (G), and cytosine (C). Each consists of a nucleotide, or a nucleotide analog. The nucleotide can be a

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In one embodiment it is preferred that at least some of said nucleotides are selected from the group consisting of nucleotide derivatives, Including deoxyribonucleic acid derivatives and ribonucleic acid derivatives.

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Coding Element Subunits and Corresponding Complementing Element Subunits

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nucleotides, nucleotide derivatives and nucleotide analogs in which one or more of a base molety and/or a phosphate molety and/or a ribose molety and/or a deoxyribose coding element subunits comprise or essentially consist of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, Including The coding element subunits are preferably selected from the group consisting of corresponding complementing element subunits capable of interacting with said moiety have been substituted by an alternative molecular entity, and the any analog or derivative thereof.

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Each nucleotide derivative can be linked to a neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond, or each nucleotide derivative can be linked to

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such as from 14 to 80, for example from 14 to 60, such as from 14 to 40, for

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The covalent chemical bond is preferably selected from the group of covalent bonds each neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond. consisting of phosphodiester bonds, phosphorothicate bonds, and peptide bonds.

Complementing Elements S

complementing efements in a linear sequence or a branched sequence. n preferably as from 3 to 10, such as from 3 to 8, for example from 3 to 6, such as from 3 to 4, for 8 to 20, for example from 8 to 15, such as from 8 to 10, for example 8, such as 9, for such as from 4 to 40, for example from 4 to 30, such as from 4 to 20, such as from 4 to 15, for example from 4 to 10, such as from 4 to 8, such as from 4 to 6, such as 4, for example from 6 to 10, such as from 6 to 8, such as 6, for example from 7 to 100, example from 8 to 60, such as from 8 to 40, for example from 8 to 30, such as from example from 3 to 80, such as from 3 to 60, such as from 3 to 40, for example from such as from 7 to 8, such as 7, for example from 8 to 100, such as from 8 to 80, for example from 10 to 100, such as from 10 to 80, for example from 10 to 60, such as 12 to 30, such as from 12 to 20, for example from 12 to 15, such as from 14 to 100, example from 2 to 60, such as from 2 to 40, for example from 2 to 30, such as from 3 to 30, such as from 3 to 20, such as from 3 to 15, for example from 3 to 15, such or example from 5 to 100, such as from 5 to 80, for example from 5 to 60, such as from 5 to 40, for example from 5 to 30, such as from 5 to 20, for example from 5 to 15, such as from 5 to 10, such as from 5 to 8, for example from 5 to 6, for example from 6 to 40, for example from 6 to 30, such as from 6 to 20, such as from 6 to 15, from 7 to 30, such as from 7 to 20, for example from 7 to 15, such as from 7 to 10, example 3, such as from 4 to 100, for example from 4 to 80, such as from 4 to 60, rom 10 to 40, for example from 10 to 30, such as from 10 to 20, for example from from 12 to 80, for example from 12 to 60, such as from 12 to 40, for example from 5, such as from 6 to 100, for example from 6 to 80, such as from 6 to 60, such as such as from 7 to 80, for example from 7 to 60, such as from 7 to 40, for example 10 to 15, such as from 10 to 12, such as 10, for example from 12 to 100, such as has a value of from 2 to 200, for example from 2 to 100, such as from 2 to 80, for 2 to 20, for example from 2 to 15, such as from 2 to 10, such as from 2 to 8, for example from 2 to 6, such as from 2 to 4, such as 2, such as from 3 to 100, for The complementing template in one embodiment preferably comprises n 9 5 ឧ 22 ജ

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from 18 to 30, such as from 18 to 20, for example from 20 to 100, such as from 20 to from 16 to 100, such as from 16 to 80, for example from 16 to 60, such as from 16 to 25 to 60, such as from 25 to 40, for example from 25 to 30, such as from 30 to 100, 40, for example from 16 to 30, such as from 16 to 20, such as from 18 to 100, such from 22 to 25, for example from 25 to 100, such as from 25 to 80, for example from example from 14 to 30, such as from 14 to 20, for example from 14 to 16, such as example from 22 to 60, such as from 22 to 40, for example from 22 to 30, such as 80, for example from 20 to 60, such as from 20 to 40, for example from 20 to 30, as from 18 to 80, for example from 18 to 60, such as from 18 to 40, for example such as from 20 to 25, for example from 22 to 100, such as from 22 to 80, for

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from 35 to 60, such as from 35 to 40, for example from 40 to 100, such as from 40 to 60 to 80, such as from 60 to 70, for example from 70 to 100, such as from 70 to 90, ' for example from 70 to 80, such as from 80 to 100, for example from 80 to 90, such example from 45 to 50, such as from 50 to 100, for example from 50 to 80, such as from 50 to 60, for example from 50 to 55, such as from 60 to 100, for example from for example from 30 to 80, such as from 30 to 60, for example from 30 to 40, such as from 30 to 35, for example from 35 to 100, such as from 35 to 80, for example 80, for example from 40 to 60, such as from 40 to 50, for example from 40 to 45, such as from 45 to 100, for example from 45 to 80, such as from 45 to 60, for as from 90 to 100.

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In some embodiments, the complementing template is attached to a solid or semi-

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The complementing template in one embodiment comprises or essentially consists of nucleotides selected from the group consisting of deoxyribonuclelc acids (DNA), ribonucleic acids (RNA), peptide nucleic acids (PNA), locked nucleic acids (LNA), and morpholinos sequences, including any analog or derivative thereof.

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essentially consist of nucleotides selected from the group consisting of DNA, RNA, In other embodiments, there is provided a complementing template comprising or PNA, LNA and morpholinos sequence, Including any analog or derivative thereof. thereof, wherein the corresponding coding elements of the template comprise or essentially consisting of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative

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template comprising a single strand of coding elements, and/or comprises a priming The complementing template is preferably amplifyable and/or comprises a single complementing elements capable of forming a double helix by hybridization to a strand of complementing elements and/or comprises a single strand of

Each complementing element is preferably linked to a neighbouring complementing element by a covalent chemical bond, or linked to each complementing element is linked to each neighbouring complementing element by a covalent chemical bond. covalent bonds consisting of phosphodiester bonds, phosphorothicate bonds, and The covalent chemical bond is in one embodiment selected from the group of peptide bonds. In other embodiments, the group of covalent bonds consist of phosphodiester bonds and phosphorothicate bonds.

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The at least one complementing element can be attached to a solld or semi-solid support.

nucleotides, including any analog or derivative thereof, amino acids, antibodies, and group consisting of nucleotides, including deoxyribonucleic acids comprising a base embodiment, it is preferred that the complementing elements are selected from the ribonucleic acids comprising a base selected from adenine (A), uracil (U), guanine derivatives, and nucleotide analogs, including any combination thereof. In one antigens, and preferably from the group consisting of nucleotides, nucleotide The complementing elements can be selected from the group consisting of selected from adenine (A), thymine (T), guanine (G), and cytosine (C), and (G), and cytosine (C). 8 22

Each nucleotide can be linked to a neighbouring nucleotide, or nucleotide analog, by neighbouring nucleotide, or nucleotide analog, by means of a covalent bond. The means of a covalent bond, Including, or each nucleotide can be linked to each covalent bond can be a phosphodiester bond or a phosphorothioate bond.

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In another embodiment, the complementing elements are natural or non-natural nucleotides selected from the group consisting of deoxyribonucleic acids and ribonucleic acids.

5 Complementing Elements and Corresponding Coding Elements

When the complementing elements are selected from the group consisting of nucleotides, nucleotide derivatives and nucleotide analogs in which one or more of a base molety and/or a phosphate molety and/or a ribose and/or a deoxyribose moiety has been substituted by an alternative molecular entity, the coding elements capable of interacting with said complementing elements comprise or essentially consist of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof.

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Each nucleotide can be linked to a neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond, or linked to each neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond. The covalent chemical bond is preferably selected from the group of covalent bonds consisting of phosphodiester bonds, phosphorothioate bonds, and peptide bonds.

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The complementing elements are in one embodiment selected from nucleotides, and the complementing elements can in one preferred embodiment be linked enzymatically by using an enzyme selected from the group consisting of template-dependent DNA- and RNA-polymerases, including reverse transcriptases, DNA-

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1gases and RNA-ligases, ribozymes and deoxyribozymes, including HIV-1 Reverse Transcriptase, AMV Reverse Transcriptase, T7 RNA polymerase, T7 RNA polymerase mutant Y639F, Sequenase, Taq DNA polymerase, Klenow Fragment (Large fragment of DNA polymerase, T7 DNA polymerase, T4 DNA Ligase, E. coli RNA polymerase, T7 DNA polymerase, Vent DNA polymerase, Pfu DNA polymerase, Vent BNA polymerase, Pfu DNA polymerase, T6 DNA polymerase, Marchites.

More preferably, the enzyme is selected from the group consisting of HIV-1 Reverse Transcriptase, AMV Reverse Transcriptase, T7 RNA polymerase, T7 RNA polymerase mutant Y639F, Sequenase, Taq DNA polymerase, Klenow Fragment

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(Large fragment of DNA polymerase I), DNA-ligase, T7 DNA polymerase, T4 DNA polymerase, and T4 DNA Ligase. The nucleotides preferably form a template or complementing template upon incorporation.

in another embodiment, the complementing elements can be selected from nucleotides, and linked by using a chemical agent, pH change, light, a catalyst, radiation, such as electromagnetic radiation, or by spontaneous coupling when being brought into reactive contact with each other.

10 Complementing Element Subunits

The complementing element preferably comprises or essentially consists of from 1 to 100 subunits, such as from 1 to 80 subunits, for example from 1 to 60 subunits,

- such as from 1 to 40 subunits, for example from 1 to 20 subunits, such as from 1 to 18 subunits, for example from 1 to 16 subunits, such as from 1 to 14 subunits, for example from 1 to 12 subunits, such as from 1 to 10 subunits, for example from 1 to 9 subunits, such as from 1 to 8 subunits, for example from 1 to 7 subunits, such as from 1 to 8 subunits, such as from 1 to 9 subunits, such as from 1 to 8 subunits, such as from 1 to 9 subunits, for example 1
- subunit, such as from 2 to 100 subunits, such as from 2 to 80 subunits, for example from 2 to 60 subunits, such as from 2 to 40 subunits, for example from 2 to 20 subunits, such as from 2 to 18 subunits, for example from 2 to 16 subunits, such as from 2 to 14 subunits, for example from 2 to 12 subunits, such as from 2 to 10 subunits, for example from 2 to 9 subunits, such as from 2 to 10 subunits, for example from 2 to 9 subunits, such as from 2 to 10 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 8 subunits, such as from 2 to 8 subunits, for example from 2 to 8 subunits, such as from 2 to 8 subunits, for example from 2 to 8 subunits, such as from 2 to 8 subunits, such
- subunits, such as from 2 to 4 subunits, for example from 2 to 5 subunits, such as from 2 to 4 subunits, for example from 2 to 3 subunits, such as 2 subunits, such as from 3 to 100 subunits, such as from 3 to 40 subunits, for example from 3 to 60 subunits, such as from 3 to 40 subunits, for example from 3 to 20 subunits, such as from 3 to 18 subunits, for example from 3 to 16 subunits, such as from 3 to 18 subunits, for example from 3 to 16 subunits, such as from 3 to 18 subunits, for example from 3 to 16 subunits, such as from 3 to 18 subunits, for example from 3 to 16 subunits, such as from 3 to 16 subunits, for example from 3 to 16 subunits, such as
- from 3 to 14 subunits, for example from 3 to 12 subunits, such as from 3 to 10 subunits, for example from 3 to 9 subunits, such as from 3 to 8 subunits, for example from 3 to 7 subunits, such as from 3 to 6 subunits, for example from 3 to 5 subunits, for example from 3 to 5 subunits, such as from 3 to 4 subunits, for example 3 subunits, for example from 4 to 100 subunits, such as from 4 to 80 subunits, for example from 4 to 20 subunits, such as from 4 to 40 subunits, for example from 4 to 20 subunits, such as from 4 to 40 subunits, for example from 4 to 20 subunits, such as from 4 to 40 subunits, for example from 4 to 20 subunits, such as from 4 to 40 subunits, for example from 4 to 50 subunits, such as from 4 to 40 subunits, for example from 6 to 80 subunits, such 80 subunit

100 subunits, such as from 6 to 80 subunits, for example from 6 to 60 subunits, such 9 subunits, such as from 6 to 8 subunits, for example from 6 to 7 subunits, such as 6 subunits, such as from 7 to 100 subunits, such as from 7 to 80 subunits, for example example from 6 to 12 subunits, such as from 6 to 10 subunits, for example from 6 to example from 8 to 60 subunits, such as from 8 to 40 subunits, for example from 8 to example from 4 to 12 subunits, such as from 4 to 10 subunits, for example from 4 to from 4 to 6 subunits, for example from 4 to 5 subunits, for example 4 subunits, such subunits, such as from 5 to 40 subunits, for example from 5 to 20 subunits, such as example from 5 to 9 subunits, such as from 5 to 8 subunits, for example from 5 to 7 such as from 9 to 40 subunits, for example from 9 to 20 subunits, such as from 9 to as from 10 to 14 subunits, for example from 10 to 12 subunits, such as 10 subunits, subunits, such as from 7 to 18 subunits, for example from 7 to 16 subunits, such as 20 subunits, such as from 8 to 18 subunits, for example from 8 to 16 subunits, such subunits, such as from 10 to 18 subunits, for example from 10 to 16 subunits, such 9 subunits, such as from 4 to 8 subunits, for example from 4 to 7 subunits, such as as from 5 to 100 subunits, such as from 5 to 80 subunits, for example from 5 to 60 subunits, for example from 7 to 9 subunits, such as from 7 to 8 subunits, such as 7 subunits, for example from 8 to 9 subunits, for example 8 subunits, such as from 9 such as from 11 to 100 subunits, such as from 11 to 80 subunits, for example from from 10 to 60 subunits, such as from 10 to 40 subunits, for example from 10 to 20 example from 9 to 12 subunits, such as from 9 to 10 subunits, such as 9 subunits, as from 8 to 14 subunits, for example from 8 to 12 subunits, such as from 8 to 10 to 100 subunits, such as from 9 to 80 subunits, for example from 9 to 60 subunits, for example from 10 to 100 subunits, such as from 10 to 80 subunits, for example 18 subunits, for example from 4 to 16 subunits, such as from 4 to 14 subunits, for subunits, such as from 5 to 6 subunits, such as 5 subunits, for example from 6 to as from 6 to 40 subunits, for example from 6 to 20 subunits, such as from 6 to 18 18 subunits, for example from 9 to 16 subunits, such as from 9 to 14 subunits, for subunits, for example from 8 to 100 subunits, such as from 8 to 80 subunits, for from 7 to 14 subunits, for example from 7 to 12 subunits, such as from 7 to 10 subunits, for example from 5 to 12 subunits, such as from 5 to 10 subunits, for subunits, for example from 6 to 16 subunits, such as from 6 to 14 subunits, for from 7 to 60 subunits, such as from 7 to 40 subunits, for example from 7 to 20 from 5 to 18 subunits, for example from 5 to 16 subunits, such as from 5 to 14 9 5 2 2 2 ဓ

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such as from 12 to 40 subunits, for example from 12 to 20 subunits, such as from 12 subunits, such as from 11 to 18 subunits, for example from 11 to 16 subunits, such as from 11 to 14 subunits, for example from 11 to 12 subunits, such as from 12 to 100 subunits, such as from 12 to 80 subunits, for example from 12 to 60 subunits,

- 80 subunits, for example from 14 to 60 subunits, such as from 14 to 40 subunits, for subunits, such as from 13 to 18 subunits, for example from 13 to 16 subunits, such as from 13 to 14 subunits, for example from 14 to 100 subunits, such as from 14 to to 18 subunits, for example from 12 to 16 subunits, such as from 12 to 14 subunits, from 13 to 60 subunits, such as from 13 to 40 subunits, for example from 13 to 20 for example from 13 to 100 subunits, such as from 13 to 80 subunits, for example Ŋ 2
- as from 17 to 40 subunits, for example from 17 to 20 subunits, such as from 17 to 18 14 to 16 subunits, such as from 15 to 100 subunits, such as from 15 to 80 subunits, subunits, such as from 17 to 80 subunits, for example from 17 to 60 subunits, such from 15 to 20 subunits, such as from 15 to 18 subunits, for example from 15 to 16 axample from 16 to 60 subunits, such as from 16 to 40 subunits, for example from example from 14 to 20 subunits, such as from 14 to 18 subunits, for example from for example from 15 to 60 subunits, such as from 15 to 40 subunits, for example subunits, such as from 16 to 100 subunits, such as from 16 to 80 subunits, for 16 to 20 subunits, such as from 16 to 18 subunits, for example from 17 to 100 ŧ
- as from 20 to 40 subunits, for example from 20 to 30 subunits, such as from 20 to 25 8 to 20 subunits, such as from 19 to 100 subunits, such as from 19 to 80 subunits, from 19 to 30 subunits, such as from 19 to 25 subunits, for example from 20 to 100 subunits, such as from 20 to 80 subunits, for example from 20 to 60 subunits, such subunits, for example from 18 to 100 subunits, such as from 18 to 80 subunits, for example from 18 to 60 subunits, such as from 18 to 40 subunits, for example from or example from 19 to 60 subunits, such as from 19 to 40 subunits, for example

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nucleotide, or a nucleotide analog. The nucleotide can be a deoxyrlbonucleic acid cytosine (C), or a ribonucleic acid comprising a base selected from adenine (A), In preferred embodiments, each subunit comprises or essentially consists of a comprising a base selected from adenine (A), thymine (T), guanine (G), and uracii (U), guanine (G), and cytosine (C).

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11 to 60 subunits, such as from 11 to 40 subunits, for example from 11 to 20

Each of said nucleotides can be linked to a neighbouring nucleotide, or nucleotide analog, by means of a covalent bond, or linked to each neighbouring nucleotide, or nucleotide analog, by means of a covalent bond. The covalent bond is preferably selected from the group consisting of phosphodiester bonds, phosphorothloate bonds, and peptide bonds.

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It is preferred in one embodiment that at least some of said nucleotides are selected from the group consisting of nucleotide derivatives, including nucleotide derivatives selected from the group consisting of deoxyribonucleic acid derivatives and ribonucleic acid derivatives.

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Complementing Element Subunits and Corresponding Coding Element
15 Subunits

When the complementing element subunits are selected from the group consisting of nucleotides, nucleotide derivatives, and nucleotide analogs in which one or more of a base moiety and/or a phosphate moiety and/or a ribose moiety and/or a deoxyribose moiety has been substituted by an atternative molecular entity, the coding element subunits capable of interacting with sald complementing element subunits preferably comprise or essentially consist of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof.

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It is preferred that each nucleotide derivative is linked to a neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond, or linked to each neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond. The covalent chemical bond can be selected from the group of covalent bonds consisting of phosphotiester bonds, phosphorothioate bonds, and peptide bonds.

Building Blocks, Cleavable Linkers and Selectively Cleavable Linkers

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in one aspect there is provided a building block comprising

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a complementing element capable of specifically recognising a coding element having a recognition group, said complementing element being selected from nucleotides, amino acids, antibodies, antigens, proteins, peptides, and molecules with nucleotide recognizing ability,

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peptides, β-peptides, γ-peptides, ω-peptides, peptides wherein the amino polyesters, polysaccharides, polycarbamates, polycarbonates, polyureas, including polyheterocyclic compounds, proteoglycans, and polysitoxanes, acid residues are in the L-form or in the D-form, vinylogous polypeptides, polysulfonamide, conjugated peptides comprising e.g. prosthetic groups, at least one functional entity selected from a precursor of α -peptides, β polypeptidylphosphonates, polyurethanes, azatides, oligo N-substituted sulfides, polynucleotides, PNAs, LNAs, morpholinos, oligo pyrrolinone, polyethylene glycols (PEG), polyethylenes, polydisulfides, polyarylene polyacetates, polystyrenes, polyvlnyl, lipids, phospholipids, glycolipids, polyoximes, polyimines, polyethyleneimlnes, polyimides, polyacetals, polycyclic compounds comprising e.g. aliphatic or aromatic cycles, peptides, γ -peptides, α -peptides, mono-, di- and tri-substituted α glycines, polyethers, ethoxyformacetal oligomers, poly-thloethers, glycopoly-peptides, polyamides, vinylogous sulfonamide peptide, ≘

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 iii) a linker or selectively cleavable linker separating the functional entity from the complementing element. The complementing element of the building block is preferably selected from a nucleotide sequence, such as a sequence of from 1 to 8 nucleotides, such as from 1 to 6 nucleotides, for example from 1 to 4 nucleotides, such as 2 nucleotides or for example 3 nucleotides.

The functional entity can be selected from a precursor of an amino acid selected from alfa amino acids, beta amino acids, gamma amino acids, di-substituted amino acids, poly-substituted amino acids, vinylogous amino acids, N-substituted glycin derivatives and other modified amino acids.

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The is also provided a composition of building blocks as defined herein, wherein at least two building blocks of the composition are different.

At least a subset of the plurality of building blocks preferably comprises one 5 complementing element and one functional entity and one linker.

In one embodiment, each building block comprises at least one reactive group type I and/or at least one reactive group type II, including one reactive group type I, two reactive groups type I, two reactive groups type I, one reactive group type II, and two reactive groups type II.

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At least one of sald reactive groups type II of the functional entity is preferably selected from the group consisting of N-carboxyanhydride (NCA), N-thiocarboxyanhydride (NTA), amine, carboxylic acid, ketone, aldehyde, hydroxyl, thiol, ester, thioester, any conjugated system of double bonds, hydrazine, N-hydroxysuccinimide ester, and epoxide.

In some embodiments, the ractive group type II is an electrophile, a nucleophile, or a

radical.

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At least a subset of said plurality of building blocks comprises a selectively cleavable linker separating the functional entity from the complementing element, wherein said selectively cleavable linker is not cleaved under conditions resulting in cleavage of cleavable linkers separating the functional entity from the complementing element of building blocks not belonging to the subset of building blocks comprising a

selectively cleavable linker. The cleavable linkers of the building blocks are cleaved without cleaving the at least one selectively cleavable linker linking the templated molecule to the complementing template, or to a complementing element, or linking said templated molecule to a templating element, or to the template that templated the synthesis of the templated molecule.

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Linkers and selectively deavable linkers can be cleaved by e.g. acid, base, a chemical agent, light, electromagnetic radiation, an enzyme, or a catalyst, with the proviso that the cleavage of the cleavable finker does result in the cleavage of the selectively cleavable linker unless this is desirable.

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In one embodiment, the length of the linker or selectively cleavable linker is in the range of from about 0.8 A to about 70 A, such as in the range of from 0.8 A to about 60 A, for example in the range of from 0.8 A to about 50 A, such as in the range of from 0.8 A to about 30 A, such as In the range of from 0.8 A to about 30 A, such

as in the range of from 0.8 A to about 25 A, for example in the range of from 0.8 A to about 20 A, such as in the range of from 0.8 A to about 18 A, for example in the range of from 0.8 A to about 16 A, such as in the range of from 0.8 A to about 14 A, for example in the range of from 0.8 A to about 12 A, such as in the range of from 0.8 A to about 10 A, for example in the range of from 0.8 A to about 8 A, such as in the range of from 0.8 A to about 8 A, such as in

the range of from 0.8 A to about 7 A, for example in the range of from 0.8 A to about 6 A, such as in the range of from 0.8 A to about 5 A, for example in the range of from 0.8 A to about 3.5 A, for example in the range of from 0.8 A to about 3.5 A, for example in the range of from 0.8 A to about 3.0 A, such as in the range of from 0.8 A to about 2.0 A, such as in the range of from 0.8 A to about 2.0 A, for example in the range of from 0.8 A to about 2.0 A, such as in the range of from 0.8 A to about 1.5 A, for example in the range of from 0.8 A to

In another embodiment, the length of the linker or selectively cleavable linker is In the range of from about 1 A to about 60 A, such as in the range of from 1 A to about 40 A for example in the range of from 1 A to about 30 A, such as in the range of

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40 A, for example in the range of from 1 A to about 30 A, such as in the range of from 1 A to about 25 A, for example in the range of from 1 A to about 20 A, such as in the range of from 1 A to about 18 A, for example in the range of from 1 A to about 16 A, such as in the range of from 1 A to about 14 A, for example in the range of from 1 A to about 14 A, for example in the range of from 1 A to about 10 A, for example

In the range of from 1 A to about 8 A, such as in the range of from 1 A to about 7 A, for example in the range of from 1 A to about 6 A, such as in the range of from 1 A to about 5 A, for example in the range of from 1 A to about 4 A, such as in the range of from 1.0 A to about 3.5 A, for example in the range of from 1.0 A to about 3.0 A, such as in the range of from 1.0 A to about 2.5 A, for example in the range of from 1.0 A to about 2.5 A, for example in the range of from

30 1.0 Å to about 2.0 Å, such as in the range of from 1.0 Å to about 1.5 Å, for example in the range of from 1.0 Å to about 1.2 Å,

In yet another embodiment, the length of the linker or selectively cleavable linker is in the range of from about 2 Å to about 40 Å, such as in the range of from 2 Å to about 30 Å, such as in the range of from 2 Å to about 30 Å, such as in the range

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of from 2 Å to about 10 Å, for example in the range of from 2 Å to about 8 Å, such as in the range of from 2 Å to about 7 Å, for example in the range of from 2 Å to about 6 about 14 A, for example in the range of from 2 A to about 12 A, such as in the range A to about 4 A, such as in the range of from 2.0 A to about 3.5 A, for example in the example in the range of from 2 A to about 16 A, such as in the range of from 2 A to A, such as in the range of from 2 A to about 5 A, for example in the range of from 2 range of from 2.0 Å to about 3.0 Å, such as in the range of from 2.0 Å to about of from 2 A to about 20 A, such as in the range of from 2 A to about 18 A, for 2.5 A, for example in the range of from 2.0 A to about 2.2 A.

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from 4 Å to about 10 Å, for example in the range of from 4 Å to about 8 Å, such as in the range of from about 4 Å to about 40 Å, such as in the range of from 4 Å to about from 4 Å to about 20 Å, such as in the range of from 4 Å to about 18 Å, for example the range of from 4 A to about 7 A, for example in the range of from 4 A to about 6 In a further embodiment, the length of the linker or selectively cleavable linker is in 30 A, such as in the range of from 4 A to about 25 A, for example in the range of 14 Å, for example in the range of from 4 Å to about 12 Å, such as in the range of in the range of from 4 Å to about 16 Å, such as in the range of from 4 Å to about A, such as in the range of from 4 A to about 5 A.

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of from 6 A to about 10 A, for example in the range of from 6 A to about 8 A, such as about 30 Å, such as in the range of from 6 Å to about 25 Å, for example in the range about 14 A, for example in the range of from 6 A to about 12 A, such as in the range In a still further embodiment, the length of the linker or selectively cleavable linker is example in the range of from 6 A to about 16 A, such as in the range of from 6 A to in the range of from about 6 Å to about 40 Å, such as in the range of from 6 Å to of from 6 A to about 20 A, such as in the range of from 6 A to about 18 A, for in the range of from 6 A to about 7 A.

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about 30 Å, such as in the range of from 8 Å to about 25 Å, for example in the range In yet another embodiment, the length of the linker or selectively cleavable linker is example in the range of from 8 Å to about 16 Å, such as in the range of from 8 Å to in the range of from about 8 Å to about 40 Å, such as in the range of from 8 Å to of from 8 A to about 20 A, such as in the range of from 8 A to about 18 A, for

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about 14 A, for example in the range of from 8 A to about 12 A, such as in the range of from 8 A to about 10 A.

Templated molecules

The templated molecules can be linked - or not linked - to the template having emplated the synthesis of the templated molecule. In one embodiment, the present invention relates to templated molecules comprising or essentially consisting of amino acids selected from the group consisting of α amino acids, β-amino acids, γ-amino acids, ∞-amino acids. 9

in various preferred embodiments the templated molecule comprises or essentially consists of one or more of natural amino acid residues, of α -amino acids, of

monosubstituted α -amino acids, disubstituted α -amino acids, monosubstituted β cetrasubstituted β-amino acids, y-amino acids, ∞-amino acids, vlnylogous amino amino acids, disubstituted B-amino acids, or trisubstituted B-amino acids, acids, and N-substituted glycines.

- have a backbone structure comprising or essentially consisting of a cyclohexane-The above-mentioned templated molecules comprising β-amino acids preferably packbone and/or a cyclopentane-backbone. ឧ
- molecules or molecular entities selected from the group of α -peptides, β -peptides, γ peptides, e-peptides, peptides wherein the amino acid residues are in the L-form or in the D-form, vinylogous polypeptides, glycopoly-peptides, polyamides, vinylogous In other embodiments, the templated molecule comprises or essentially consists of peptides, α -peptides, mono-, dl- and trl-substituted α -peptides, β -peptides, γ sulfonamide peptide, polysulfonamide, conjugated peptides comprising e.g. 22
 - polyureas, polypeptidylphosphonates, polyurethanes, azatides, oligo N-substituted glycols (PEG), polyethylenes, polydisulfides, polyarylene sulfides, polynucleotides, prosthetic groups, polyesters, polysaccharides, polycarbamates, polycarbonates, glycines, polyethers, ethoxyformacetal oligomers, poly-thioethers, polyethylene PNAs, LNAs, morpholinos, oligo pyrrolinone, polyoximes, polyimines, ಜ
- polyethyleneimines, polyimides, polyacetals, polyacetates, polystyrenes, polyvinyl, 33

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lipids, phospholipids, glycolipids, polycyclic compounds comprising e.g. aliphatic or aromatic cycles, including polyheterocyclic compounds, proteoglycans, and polysiloxanes, inlcuding any combination thereof.

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Neighbouring residues of the templated molecules according to the invention can be linked by a chemical bond selected from the group of chemical bonds consisting of conds, carbonate bonds, urea bonds, phosphonate bonds, urethane bonds, azatide bonds, peptoid bonds, ether bonds, ethoxy bonds, thioether bonds, single carbon bonds, double carbon bonds, triple carbon bonds, disulfide bonds, sulfide bonds, peptide bonds, sulfonamide bonds, ester bonds, saccharide bonds, carbamate phosphodiester bonds, oxime bonds, imine bonds, imide bonds, including any combination thereof.

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NHCH2CHRCO-;-NHCHRCH2CO-;-COCH2-;-COS-;-CONR-;-COO-;-CSNH-;can in one aspect comprise or essentially consist of a molecular group selected from Also, the backbone structure of the templated molecules according to the invention NHN(R)CO-; -NHB(R)CO-; -NHC(RR')CO-; -NHC(=CHR)CO-; -NHC, H, CO-; -CH2 NH-; -CH2CH2-; -CH2 S-; -CH2 SO-; -CH2SO2-; -CH(CH3)S-; -CH=CH-; -NHCO.; -NHCONH-; -CONHO.; -C(=CH2)CH2-; -PO2NH-; -PO2 CH2-; -PO2 CH2N*-; -SO2NH-; and lactams, Including any combination thereof.

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In other embodiments of the invention, the templated molecules are not of polymeric

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precursors consisting of a-amino acid precursors, β-amino acid precursors, y-amino The precursor is in one embodiment preferably selected from the group of acid precursors, and e-amino acid precursors. 22

comprising at least one repetitive sequence of functional groups, such as at least templated molecules also includes molecules wherein any sequence of at least three functional groups repeated at least twice in the templated molecule. The In some embodiment, the templated molecule is an oligomer or a polymer three functional groups occurs only once.

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Some preferred templated molecules preferably comprise or essentially consist of at least 2 different functional groups, such as at least 3 different functional groups, for example at least 4 different functional groups, such as at least 5 different functional

- groups, for example at least 6 different functional groups, such as at least 7 different functional groups, for example at least 8 different functional groups, such as at least such as more than 10 different functional groups. The functional groups can also be 9 different functional groups, for example at least 10 different functional groups, dentical, ιO
- rom 2 to 200, for example from 2 to 100, such as from 2 to 80, for example from 2 sach comprising at least one residue, wherein the plurality of residues is preferably comprising a polymer comprising a plurality of covalently linked functional groups In one preferred aspect of the invention there is provided a templated molecule to 60, such as from 2 to 40, for example from 2 to 30, such as from 2 to 20, for 2
- example from 2 to 15, such as from 2 to 10, such as from 2 to 8, for example from 2 to 80, such as from 3 to 60, such as from 3 to 40, for example from 3 to 30, such as to 6, such as from 2 to 4, for example 2, such as from 3 to 100, for example from 3 rom 3 to 20, such as from 3 to 15, for example from 3 to 15, such as from 3 to 10, such as from 3 to 8, for example from 3 to 6, such as from 3 to 4, for example 3, \$
- example from 5 to 100, such as from 5 to 80, for example from 5 to 60, such as from uch as from 4 to 100, for example from 4 to 80, such as from 4 to 60, such as from example from 4 to 10, such as from 4 to 8, such as from 4 to 6, for example 4, for 5 to 40, for example from 5 to 30, such as from 5 to 20, for example from 5 to 15, 4 to 40, for example from 4 to 30, such as from 4 to 20, such as from 4 to 15, for 8
- such as from 6 to 100, for example from 6 to 80, such as from 6 to 60, such as from such as from 5 to 10, such as from 5 to 8, for example from 5 to 6, for example 5, such as from 7 to 80, for example from 7 to 60, such as from 7 to 40, for example 6 to 40, for example from 6 to 30, such as from 6 to 20, such as from 6 to 15, for example from 6 to 10, such as from 6 to 8, such as 6, for example from 7 to 100, 22
- such as from 7 to 8, for example 7, for example from 8 to 100, such as from 8 to 80, from 8 to 20, for example from 8 to 15, such as from 8 to 10, such as 8, for example from 7 to 30, such as from 7 to 20, for example from 7 to 15, such as from 7 to 10, or example from 8 to 60, such as from 8 to 40, for example from 8 to 30, such as 9, for example from 10 to 100, such as from 10 to 80, for example from 10 to 60, ജ
 - such as from 10 to 40, for example from 10 to 30, such as from 10 to 20, for 32

example from 10 to 15, such as from 10 to 12, such as 10, for example from 12 to

100, such as from 12 to 80, for example from 12 to 60, such as from 12 to 40, for

from 14 to 100, such as from 14 to 80, for example from 14 to 60, such as from 14 to

example from 12 to 30, such as from 12 to 20, for example from 12 to 15, such as

from 16 to 40, for example from 16 to 30, such as from 16 to 20, such as from 18 to

such as from 16 to 100, such as from 16 to 80, for example from 16 to 60, such as

40, for example from 14 to 30, such as from 14 to 20, for example from 14 to 16,

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example from 18 to 30, such as from 18 to 20, for example from 20 to 100, such as

100, such as from 18 to 80, for example from 18 to 60, such as from 18 to 40, for

20 to 30, such as from 20 to 25, for example from 22 to 100, such as from 22 to 80,

for example from 22 to 60, such as from 22 to 40, for example from 22 to 30, such

from 20 to 80, for example from 20 to 60, such as from 20 to 40, for example from

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polyurethanes, azatides, oligo N-substituted glycines, polyethers, ethoxyformacetal polydisulfides, polyarylenė sulfides, polynudeotides, PNAs, LNAs, morpholinos, polycarbamates, polycarbonates, polyureas, polypeptidylphosphonates, oligomers, poly-thioethers, polyethylene glycols (PEG), polyethylenes,

- polyacetals, polyacetates, polystyrenes, polyvinyl, lipids, phospholipids, glycolipids, plurality of residues is preferably from 2 to 200, for example from 2 to 100, such as polyheterocyclic compounds, proteoglycans, and polysiloxanes, and wherein the polycyclic compounds comprising e.g. aliphatic or aromatic cycles, including oligo pyrrolinones, polyoximes, polyimines, polyethyleneimines, polyimides, ß
- rom 3 to 100, for example from 3 to 80, such as from 3 to 60, such as from 3 to 40, from 2 to 80, for example from 2 to 60, such as from 2 to 40, for example from 2 to 30, such as from 2 to 20, for example from 2 to 15, such as from 2 to 10, such as for example from 3 to 30, such as from 3 to 20, such as from 3 to 15, for example from 2 to 8, for example from 2 to 6, such as from 2 to 4, for example 2, such as 2
 - such as from 4 to 60, such as from 4 to 40, for example from 4 to 30, such as from 4 such as from 3 to 4, for example 3, such as from 4 to 100, for example from 4 to 80, to 20, such as from 4 to 15, for example from 4 to 10, such as from 4 to 8, such as from 3 to 15, such as from 3 to 10, such as from 3 to 8, for example from 3 to 6, from 4 to 6, for example 4, for example from 5 to 100, such as from 5 to 80, for 5
- example from 5 to 6, for example 5, such as from 6 to 100, for example from 6 to 80, such as from 6 to 60, such as from 6 to 40, for example from 6 to 30, such as from 6 example from 5 to 60, such as from 5 to 40, for example from 5 to 30, such as from to 20, such as from 6 to 15, for example from 6 to 10, such as from 6 to 8, such as 5 to 20, for example from 5 to 15, such as from 5 to 10, such as from 5 to 8, for ន

from 60 to 80, such as from 60 to 70, for example from 70 to 100, such as from 70 to

90, for example from 70 to 80, such as from 80 to 100, for example from 80 to 90,

such as from 90 to 100.

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as from 50 to 60, for example from 50 to 55, such as from 60 to 100, for example

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from 25 to 60, such as from 25 to 40, for example from 25 to 30, such as from 30 to

as from 22 to 25, for example from 25 to 100, such as from 25 to 80, for example

100, for example from 30 to 80, such as from 30 to 60, for example from 30 to 40,

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such as from 30 to 35, for example from 35 to 100, such as from 35 to 80, for

example from 35 to 60, such as from 35 to 40, for example from 40 to 100, such as

from 40 to 80, for example from 40 to 60, such as from 40 to 50, for example from

10 to 45, such as from 45 to 100, for example from 45 to 80, such as from 45 to 60, for example from 45 to 50, such as from 50 to 100, for example from 50 to 80, such

- as from 7 to 40, for example from 7 to 30, such as from 7 to 20, for example from 7 example from 8 to 30, such as from 8 to 20, for example from 8 to 15, such as from to 15, such as from 7 to 10, such as from 7 to 8, for example 7, for example from 8 6, for example from 7 to 100, such as from 7 to 80, for example from 7 to 60, such to 100, such as from 8 to 80, for example from 8 to 60, such as from 8 to 40, for 33
 - 10, for example from 12 to 100, such as from 12 to 80, for example from 12 to 60, 8 to 10, such as 8, for example 9, for example from 10 to 100, such as from 10 to such as from 10 to 20, for example from 10 to 15, such as from 10 to 12, such as 80, for example from 10 to 60, such as from 10 to 40, for example from 10 to 30, such as from 12 to 40, for example from 12 to 30, such as from 12 to 20, for ဓ

portions, at least one portion selected from the group of polymer portlons consisting

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generating a polymer comprising, exclusively or in combination with additional

of α -peptides, β -peptides, γ -peptides, α -peptides, mono-, di- and tri-substituted α -

In another preferred aspect of the invention there is provided a templated molecule

comprising a polymer comprising a plurality of covalently linked functional groups

each comprising a residue, wherein the covalently linked residues are capable of

example from 12 to 15, such as from 14 to 100, such as from 14 to 80, for example

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conjugated peptides comprising e.g. prosthetic groups, polyesters, polysaccharides,

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residues are in the L-form or in the D-form, vinylogous polypeptides, glycopoly-

peptides, polyamides, vinylogous sulfonamide peptides, polysulfonamides,

peptides, β-peptides, γ-peptides, ω-peptides, peptides wherein the amino acid

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from 70 to 100, such as from 70 to 90, for example from 70 to 80, such as from 80 to from 25 to 30, such as from 30 to 100, for example from 30 to 80, such as from 30 to from 16 to 20, such as from 18 to 100, such as from 18 to 80, for example from 18 to for example from 22 to 30, such as from 22 to 25, for example from 25 to 100, such from 14 to 60, such as from 14 to 40, for example from 14 to 30, such as from 14 to 22 to 100, such as from 22 to 80, for example from 22 to 60, such as from 22 to 40, example from 40 to 100, such as from 40 to 80, for example from 40 to 60, such as from 40 to 50, for example from 40 to 45, such as from 45 to 100, for example from example from 20 to 100, such as from 20 to 80, for example from 20 to 60, such as 45 to 80, such as from 45 to 60, for example from 45 to 50, such as from 50 to 100, for example from 50 to 80, such as from 50 to 60, for example from 50 to 55, such from 20 to 40, for example from 20 to 30, such as from 20 to 25, for example from example from 16 to 60, such as from 16 to 40, for example from 16 to 30, such as 60, for example from 30 to 40, such as from 30 to 35, for example from 35 to 100, 20, for example from 14 to 16, such as from 16 to 100, such as from 16 to 80, for 60, such as from 18 to 40, for example from 18 to 30, such as from 18 to 20, for as from 25 to 80, for example from 25 to 60, such as from 25 to 40, for example as from 60 to 100, for example from 60 to 80, such as from 60 to 70, for example such as from 35 to 80, for example from 35 to 60, such as from 35 to 40, for 100, for example from 80 to 90, such as from 90 to 100.

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The templated molecule in one embodiment is preferably one, wherein the covalently linked residues are capable of generating a polymer comprising, exclusively or in combination with additional portions selected from the group, at least one portion selected from the group of polymer portions consisting of α-peptides, β-peptides, γ-peptides, α-peptides, mono-, di- and tri-substituted α-peptides, β-peptides, γ-peptides, α-peptides wherein the amino acid residues are in the L-form or in the D-form, and vinylogous polypeptides.

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In one particular embodiment, the templated molecule is one wherein the covalently linked residues are capable of generating a polysaccharaide.

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In another aspect there is provided a templated molecule comprising a sequence of functional groups, wherein neighbouring functional groups are linked by a molecular moiety that is not natively associated with said functional groups.

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Additional aspect of the present invention relates to i) a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule does not comprise or consist of an α-peptide or a nucleotide, ii) a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule does not comprise or consist of a monosubstituted α-peptide or a nucleotide, and iii) a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule does not

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Compositions of Templated Molecules

comprise or consist of a peptide or a nucleotide.

templated molecules, such as more than or about 104 different templated molecules, than or about 10⁶ different templated molecules, for example more than or about 10⁷ as more than or about 1010 different templated molecules, for example more than or herein immediately above, can be present in a composition of templated molecules, molecules, for example more than or about 108 different templated molecules, such about 1011 different templated molecules, such as more than or about 1012 different example more than or about 10¹⁵ different templated molecules, such as more than different templated molecules, such as more than or about 1018 different templated wherein said composition comprises a plurality of more than or about 103 different different templated molecules, such as more than or about 108 different templated or about 1016 different templated molecules, for example more than or about 1017 for example more than or about 105 different tempfated molecules, such as more The templated molecules according to the invention, including those mentioned molecules, such as more than or about 10¹⁴ different templated molecules, for templated molecules, for example more than or about 1013 different templated molecules. 5 8 22

The composition in some embodiments preferably further comprises the template capable of templating each templated molecule, or a subset thereof. Accordingly, in one preferred aspect of the present invention, there is provided i) a composition comprising a templated molecule and the template capable of templating the templated molecule, or II) a composition comprising a templated molecule and the templated molecule and the templated molecule.

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Various preferred features of the templated molecules either i) linked to the template capable of templating the synthesis of the templated molecule, or ii) present in a composition further comprising the template capable of templating the synthesis of the templated molecule is listed herein immediately below.

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When being present in such compositions, it is preferred that i) the template does not consist exclusively of natural nucleotides, when the templated molecule is a peptide comprising exclusively monosubstituted α-amino acids, ii) the template is not a natural nucleotide, when the templated molecule is a natural α-peptide, iii) the template is not a nucleotide, when the templated molecule is a natural α-peptide, iv) the template is not a nucleotide, when the templated molecule is a monosubstituted α-peptide, v) the template is not a nucleotide, when the templated molecule is a peptide, vI) the template is not a natural nucleotide, when the templated molecule is a peptide, and vii) the template is not a nucleotide, when the templated molecule is

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20 Templated Molecules Linked to the Template that Templated the Synthesis of the Templated Molecule

in one preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the templated molecule does not comprise or consist of an α-peptide

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In another preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the templated molecule does not comprise a monosubstituted α-peptide.

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In yet another preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule,

 $5\,$ wherein the templated molecule does not comprise or consist of an $\alpha\text{-peptide}$ or a nucleotide.

In a still further aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template is not a natural nucleotide, when the templated molecule is an α -peptide.

In a still further preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template does not consist exclusively of natural nucleotides, when the templated molecule is a peptide comprising exclusively monosubstituted α-amino acids.

In a still further preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template is not a natural nucleotide, when the templated molecule is a

natural α-peptide.

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In an even further preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template is not a nucleotide, when the templated molecule is a natural α-peptide.

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In a still further preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or femplate that templated the synthesis of the templated molecule, wherein the templated so not a nucleotide, when the templated molecule is a

monosubstituted α-peptide.

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In a still further preferred aspect of the present Invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or templated molecule, wherein the template that templated the synthesis of the templated molecule, wherein the template is not a nucleotide, when the templated molecule is an α-peptide.

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In a still further preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template is not a natural nucleotide, when the templated molecule is a peptide.

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In a still further preferred aspect of the present invention there is provided a templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template Is not a nucleotide, when the templated molecule is a peptide.

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The templated molecule can be obtained according to the methods described herein

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In even further aspects there is provided

- a templated molecule comprising a sequence of covalently linked building blocks;
- ii) a templated molecule comprising a sequence of covalently linked building blocks, wherein the sequence of covalently linked building blocks comprises

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a sequence of complementing elements forming a complementing template capable of complementing the template that templated the synthesis of the templated molecule, and wherein the templated molecule is linked to the complementing template or template that templated its synthesis; and

a templated molecule according to any of the previous claims, wherein the
templated molecule comprises a sequence of functional entitles comprising
at least one functional group, and optionally at least one reactive group type
II, and wherein each functional entity is linked to a complementing element
or a template that templated the synthesis of the templated molecule.

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Uses of Templated Molecules

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The templated molecules according to the present invention can be used for a variety of commercial pruposes.

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In one aspect, there is provided a method for screening templated molecules potentially having a predetermined activity, said method comprising the step of providing a target molecule or a target entity, including a surface, and obtaining templated molecules having an affinity for - or an effect on - said target molecule or

target entity.

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Another aspect relates to a method for assaying an activity potentially associated with a templated molecules, said method comprising the step of providing a target molecule or a target entity, including a surface, and obtaining templated molecules.

25 having an affinity for - or an effect on - said target molecule or target entity, and determining the activity of the templated molecule.

molecules having a predetermined activity, said method comprising the step of performing a selection procedure and selecting templated molecules based on predetermined selection criteria.

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Yet another aspect provides a method for selecting complexes or templated

There is also provided a method for screening a composition of molecules having a predetermined activity comprising:

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 sstablishing a first composition of templated molecules as described herein, or produced as defined herein by any method for preparing templated molecules, exposing the first composition to conditions enriching said first composition with templated molecules having the predetermined activity, and

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 iii) optionally amplifying the templated molecules of the enriched composition obtaining a second composition,

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- iv) further optionally repeating step ii) to iii), and
- obtaining a further composition having a higher ratio of templated molecules having the specific predetermined activity.

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In one embodiment, the method further comprises a step of mutating the templated molecules, wherein sald mutagenesis can take place prior to carrying out step iii), simultaneously with carrying out step iii), or after carrying out step iii). The mutagenesis can be carried out as random or site-directed mutagenesis.

Step iii) of the method preferably comprises a 10¹ to 10¹⁶-fold amplification, and steps ii) and iii) can be repeated, such as at least 2 times, 3 times, 5 times, or at

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least 10 times, such as at least 15 times.

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The method can comprise a further step of identifying the templated molecule having the predetermined activity, and sald identification can be conducted e.g. by analysing the template and/or complementary template physically or by other means associated with the molecule.

The conditions enriching the first composition can comprise the further providing a binding partner to said templated molecule having the predetermined activity, wherein said binding partner is directly or indirectly immobilised on a support.

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The conditions enriching the composition can involve any state of the art method, including any one or more of electrophoretic separation, gelfiltration, immunoprecipitation, isoelectric focusing, centrifugation, and immobilization. The conditions enriching the composition can also comprise the further step of providing cells capable of internalising the templated molecule, or performing any interaction with the templated molecule having the predetermined activity.

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The predetermined activity of the templated molecule is preferably an enzymatic activity or a catalytic activity.

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In another aspect there is provided a method for amplifying the complementing template or the template that templated the synthesis of the templated molecule having, or potentially having a predetermined activity, said method comprising the

step of contacting the template with amplification means, and amplifying the template. The method for amplifying the complementing template or the template that templated the synthesis of the templated molecule having, or potentially having, a predetermined activity, preferably comprises the steps of I) contacting the template with amplification means, and amplifying the template, and ii) obtaining the templated molecule in an at least two-fold increased amount.

In another aspect there is provided a method for altering the sequence of a templated molecule, including generating a templated molecule comprising a novel or altered sequence of functional groups, wherein said method preferably comprises the steps of

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 providing a first complementing template or a first template capable of templating the first templated molecule, or a plurality of such first complementing templates or first templates capable of templating a plurality of first templated molecules,

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ii) mutating or modifying the sequence of the first complementing template
or the first template, or the plurality of first complementing templates or
first templates, and generating a second template or a second complementing template, or a plurality of second templates or second complementing templates,

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wherein said second template(s) or complementing template(s) is capable of templating the synthesis of a second templated molecule, or a plurality of second templated molecules,

wherein said second templated motecule(s) comprises a sequence of covalently linked, functional groups that is not identical to the sequence of functional groups of the first templated molecule(s), and optionally

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 iii) templating by means of said second template(s) or complementing template(s) a second templated molecule, or a plurality of such second templated molecules.

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In yet another aspect there is provided a method for altering the sequence of a templated molecule, including generating a templated molecule comprising a novel or altered sequence of functional groups, wherein said method preferably comprises the steps of

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 i) providing a plurality of first complementing templates or first templates capable of templating a plurality of first templated molecules,

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 ii) recombining the sequences of the plurality of first complementing templates or first templates, and generating a second template or a second complementing template, or a plurality of second templates or second complementing templates,

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wherein said second template(s) or complementing template(s) is capable of templating the synthesis of a second templated molecule, or a plurality of second templated molecules.

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wherein said second templated molecule(s) comprises a sequence of covalently linked, functional groups that is not identical to the sequence of functional groups of the first templated molecule(s), and optionally

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 iii) templating by means of said second template(s) or complementing template(s) a second templated molecule, or a plurality of such second templated molecules.

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The methods can preferably comprise the further step of amplifying the complementing template or the template that templated the synthesis of the templated molecule, wherein said amplification step taking place prior to, simultaneously with, or after the step of mutagenesis or recombination.

When mutagenesis is used, it can be used as either site-directed mutagenesis, cassette mutagenesis, chemical mutagenesis, unique site-elimination (USE), error-prone PCR, error-prone DNA shuffling. Mutagenesis preferably involves DNA shuffling and/or any form of recombination including homologous recombination either in vivo or in vitro.

Variants and functional equivalents of templated molecules

The present invention is also directed to any variant and functional equivalent of a templated molecule. The variants and functional equivalents may be obtained by any state-of-the-art-method for modifying templated molecules in the form of polymers, including peptides.

In the context of the templated molecules of the present invention, molecules are said to be homologous if they contain similar backbone structures and/or similar functional groups. Functional groups, or molecular entities of functional groups, are divided into three homology groups. The charged functional groups, the hydrophobic groups, and the hydrophilic groups. When a functional group includes two or three molecular entities belonging to different homology groups, the functional group is said to belong to the two or three different homology groups.

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Homology is measured in percent (%). As an example, the sequences AABBCA-CAAA and BBAACACBBB (where A, B and C denotes a functional group belonging to homology group A, B, and C, respectively) are 30 percent homologous.

Example 1 to 7: Preparation of the mononucleotide building block (I)

Building block I may be prepared according to the general scheme shown below: 2

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Example 1: Preparation of 3-tert-Butoxycarbonylamino-propionic acid (N-

Boc-6-alanine)(1a)

To a solution of P-alanine (2,25 g, 25 mmol) in aq. NaHCO₃ (25 mL) were added ditert-butyl dicarbonate (4,36 g, 20 mmol) and acetonitrile (25 mL). The reaction mixture was stirred at room temperature for 18 h.

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The product was extracted into EtOAc (3 x 50 mL), dried (Na₂SO₄), and evaporated EtOAc (100 mL) was added and pH was adjusted to 4-5 by addition of NaH2PO4. to dryness under vacuum to afford 3.71 g (98%) 14 NMR (CDCl₃) § 11 (14, br s, COOH), 5,07 (14, br s, NH), 3,40 (2H, m), 2,58 (2H, m), 1,44 (9H, s, 'Bu). 5

Example 2: Preparation of N-Boc-B-alanine propargyl ester(1b).

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dissolved in EtOAc (25 mL). Dicyclohexyl-carbodilmide (DCC, 2.06 g, 10 mmol) was N-Boc-β-alanine (1,91 g, 10.1 mmol) and propargyl alcohol (0.675 g, 12 mmol) were mixture was filtered and evaporated to dryness under vacuum. Crude product yield added to the solution and after 16 h of stirring at room temperature, the reaction

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Example 3: Preparation of 5-lodo-2'-deoxyuridine 3',5'-Di-tertbutyldimethylsilyl Ether(1c).

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mmol) was dissolved in anhydrous DMF (10 mL). A solution of tert-butyldimethylsilyl chloride (2.24 g, 14.9 mmol) in anhydrous DMF (5 mL) was added and the resulting 5-lodo-2'-deoxyuridine (Aldrich, 2.39 g, 6.7 mmol) and imidazole (2.025 g, 29.7 mixture was stirred for 16 h at room temperature.

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removed under reduced pressure to leave a colourless oil that solidified on standing. The reaction mixture was poured into EtOAc (400 mL), washed with NH,CI (50% sat. aq, 80 mL) followed by water (80 mL). After drying with Na₂SO₄, EtOAc was Recrystallization in n-hexane (14 mL) afforded 2.64 g, 80%.

s, 'Bu); 0.90(9H, s, 'Bu); 0.15 (3H, s, CH₃); 0.13 (3H, s, CH₃); 0.08 (3H, s, CH₃); 0.07 4.05 (1H, dd); 3.92 (1H, dd); 3.78 (1H, dd); 2,32 (1H, ddd); 2.05 (1H, ddd); 0.95(9H, 'H NMR (CDCl₃) 8 8.18 (1H, br s, NH); 8.10 (1H, s); 6,23 (1H, dd); 4,40 (1H, dt); 5

Example 4: Preparation of compound (1d)

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Compound (1d)

8.9 mmol) and triethylamine (0.585 g. 5.8 mmol) in 10 mL dry DMF were stirred at A solution of iodo silyl ether (1c) (1.62 g, 2.7 mmol), N-Boc-β-alanine(1a) (2.03 g, room temperature. N2 was passed through the solution for 20 min. Tetrakis(triphenylphosphine)palladium(0) (269 mg, 0.2 mmol) and copper(I) iodide (90 mg, 0.4 mmol) were added and the reaction mixture was stirred at room temperature for 32 h.

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EtOAc (100 mL) was poured into the reaction mixture, followed by washing (aq Na-HCO₃ (50 mL); brine (50 mL)), drying (Na₂SO₄), and removal of solvent by vacuum evaporation. The crude product (2.4 g) was purified by silica column chromatography eluting with EtOAc:Heptane gradient (1:2)(5:3) (v/v). Product yield 1.15 g, 60% 2

1'-H), 4.82 (2H, s, CH₂O), 4,39 (1H, m, 3'-H), 3.97 (1H, m, 4'-H), 3.80 (2H, dd, 5',5''- . H), 3.40 (2H, m, CH2N), 2.58 (2H, t, CH2), 2.2 (1H, m, 2'-H), 2.0 (1H, m, 2"-H), 1.45 (9H, s, 'Bu), 0.93 (9H, s, 'Bu), 0.89 (9H, s, 'Bu), 0.15 (3H, s, CH₃), 0.13 (3H, s, CH₃), H NMR (CDCI₃) § 8.45 (1H, s), 8.05 (1H, s, 6-H), 7.35 (1H, bs, NH), 6.25 (1H, dd, 0.08 (3H, s, CH₃), 0.07 (3H, s, CH₃).

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Example 5: Preparation of compound (1e) ឧ

The reaction mixture was evaporated and purified by silica column chromatography eluting with dichloromethane(DCM):methanol(MeOH) gradient (95:5)-(88:12) (v/v). acid (75 mg, 1.25 mmol) and tetrabutylammonium fluoride trihydrate (TBAF) (189 A solution of N-Boc-β-alanine silyl ether (1d) (100 mg, 0.15 mmol), glacial acetic mg, 0.6 mmol) in 2 mL dry THF was stirred at room temperature for 3 d. Product yield 26 mg, 38%. 22 ജ

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H NMR (CD3OD) 8 8.35 (1H, s, 6-H), 6.15 (1H, t, 1'-H), 4.80 (2H, s, CH2O), 4,32 (1H, dt, 3-H), 3.86 (1H, q, 4'-H), 3.70 (2H, dd, 5',5"-H), 3.24 (2H, m, CH₂N), 2.47 (2H, t, CH₂), 2,28-2.10 (1H, m, 2',2"-H), 1.44 (9H, s, 'Bu).

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Example 6: Preparation of compound (1f)

COMPOUND 1f 9

(POCI₃) in dry trimethylphosphate was added (100 µL stock solution (104 mg/mL), N-Boc-β-alanine nucleoside (1e) (26 mg, 57μmol) was dissolved in 200 μL dry trimethylphosphate. After cooling to 0 °C, a solution of phosphorus oxychloride 68 μmol). The reaction mixture was stirred at 0 °C for 2h.

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Subsequently a solution of tributylammonium pyrophosphate (Sigma P-8533) (67.8 mg, 143 µmol in 300 µL dry DMF) and tributylamine (26.9 mg, 145 µmol in 150 µL dry DMF) was added at 0 °C. The reaction was stirred at room temperature for 3 min. and then stopped by addition of 1 mL 1.0 M triethylammonium hydrogencar-

Example 7: Preparation of compound 1

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COMPOUND I

Removal of N-Boc protection group.

to pH = 1 using HCI and incubated at room temperature for 30 minutes. The mixture Following phosphorylation, 50 µl of the phosphorylation reaction mixture is adjusted is adjusted to pH 5.5 using equimolar NaOH and Na-acetate (pH 5.5) before purification on TLC.

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Purification of nucleotide derivatives using thin-layer chromatography (TLC)

tration of each nucleotide derivative was evaluated by UV-absorption prior to use in was resuspended in 50-100 µl H₂O to a final concentration of 1-3 mM. The concenfrom the nucleotides derivatives using 100% methanol as running solution. Subseshadowing. Klesel containing the nucleotide-derivative was isolated and extracted From the crude mixture, 20 samples of 2 µl were spotted on kieselgel 60 F₂₅₄ TLC quently, the TLC plate is air-dried and the nucleotide-derivative identified by UVcentrifugation and the supernatant was dried in vacuo. The nucleotide derivative (Merck). Organic solvents and non-phosphorylated nucleosides were separated twice using 10 mM Na-acetate (pH = 5.5) as solvent. Kieselgel was removed by polymerase extension reactions.

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Examples 8 to 13: Preparation of the mononucleotide building block (ii)

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Building block It may be prepared according to the general scheme shown below:

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Example 8: Preparation of N-Boc-3-phenyl-8-alanine (2a).

COMPOUND 2a

To a solution of 3-amino-3-phenylpropionic acid (3.30 g, 20 mmol) in NaHCO₂ (50% sat. aq, 25 mL) were added di-fert-butyl dicarbonate (4,36 g, 20 mmol) and acetonitie (30 mL). The reaction mixture was stirred at room temperature for 18 h. Di-fert-butyl dicarbonate (4,36 g, 20 mmol) was added and the reaction mixture was stirred at room temperature for 18 h.

EtOAc (100 mL) was added and pH was adjusted to 4-5 by addition of NaH₂PO₄. The product was extracted into EtOAc (3 x 100 mL), dried (Na₂SO₄), and evaporated to dryness under vacuum to afford crude product 5.6 g (105%)

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Example 9: Preparation of 5-(3-Hydroxypropyn-1-yl)-2'-deoxyuridine 3',5'-Ditert-butyldimethylslyl Ether(2b).

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COMPOUND 2b

A solution of lodo sily ether (3) (1.30 g, 2.2 mmol), propargyl alcohol (0.386 g, 6.9 mmol) and triethylamine (0.438 g, 4.3 mmol) in 7 mL dry DMF was deaeraed with N₂. Tetrakis(triphenylphosphine)palladium(0) (228 mg, 0.2 mmol) and copper(l) iodide (120 mg, 0.4 mmol) were added and the reaction mixture was stirred at room temperature for 32 h.

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EtOAc (100 mL) was poured into the reaction mixture, followed by washing (aq Na-HCO₃ (50 mL); brine (50 mL)), drying (Na₂SO₄), and removal of solvent by vacuum evaporation.

The crude product (1.73 g) was purified by silica column chromatography eluting 5 with EtoAc:Heptane gradient (2.3)-(3.2) (v/v). Product yield 0.713 g, 63%.

¹H NMR (CDCl₃) 8 8.47 (1H, s), 8.05 (1H, s, 6-H), 6.29 (1H, dd, 1'-H), 4,42 (2H, s, CH₂), 4,39 (1H, m, 3'-H), 3.98 (1H, m, 4'-H), 3.83 (2H, dd, 5',5"-H), 2,32 (1H, m, 2'-H), 0.93 (9H, s, Bu), 0.89 (9H, s, Bu), 0.15 (3H, s, CH₃), 0.13 (3H, s, CH₃), 0.08 (3H, s, CH₃), 0.07 (3H, s, CH₃).

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Example 10: Preparation of compound (2c)

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COMPOUND 2c

A-Boc-3-phenyl-β-alanine (8)(265 mg, 1.0 mmol) and compound (2b) (255 mg, 0.5 mmol) were dissolved in THF (15 mL). Diisopropyl-carbodiimide (DIC, 126 mg, 1 mmol) and 4-dimethylaminopyridin (DMAP, 10 mg) were added to the solution, and after 16 h of stirring at room temperature the reaction mixture was poured into EtOAc (100 mL), washed with NaHCO₃ (50% sat. aq, 50 mL), dried (Na₂SO₄), filtered and evaporated under vacuum.

25 The crude product was purified by silica column chromatography eluting with ErOAc:Heptane gradient (1.2)-(2.3) (v/v). Product yield 335 mg, 88%. 'H NMR (CDCI₃) δ 8.49 (1H, s), 8.04 (1H, s, 6-H), 7.29 (5H, m, Ph), 6.27 (1H, dd, 1'-H), 5.5 (1H, bd), 5.09 (1H, m), 4,80 (2H, s, CH₂), 4,39 (1H, m, 3'-H), 3.98 (1H, m, 4'-

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(9H, s, 'Bu), 0.91 (9H, s, 'Bu), 0.89 (9H, s, 'Bu), 0.15 (3H, s, CH₃), 0.13 (3H, s, CH₃), H), 3.82 (2H, dd, 5',5"-H), 2,87 (2H, d), 2.29 (1H, m, 2'-H), 2.01 (1H, m, 2"-H), 1.41 0.08 (3H, s, CH₃), 0.07 (3H, s, CH₃).

Example 11: Preparation of compound 2d

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COMPOUND 2d 9

mmol) and tetrabutylammonium fluoride trihydrate (TBAF) (500 mg, 1.58 mmol) in 6 A solution of compound (2c) (334 mg, 440 µmol), glacial acetic acid (190 mg, 3.15 mL dry THF was stirred at room temperature for 18 h. The reaction mixture was evaporated and purified by silica column chromatography eluting with (DCM):(MeOH) gradient (95:5)-(9:1) (v/v). Product yield 122 mg, 52%.

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'H NMR (CDC13) 8 10.1 (1H, s), 8.24 (1H, s, 6-H), 7.3 (5H, m, Ph), 6.37 (1H, dd, 1'-H), 5.6 (1H, bs), 5.09 (1H,m), 4,79 (2H, s, CH₂), 4,52 (1H, m, 3'-H), 4.0 (1H, m, 4'-H), 3.85 (2H, dd, 5',5"+H), 2,87 (2H, d), 2.4 (1H, m, 2'-H), 2.25 (1H, m, 2"-H), 1.4

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Example 12: Preparation of compound (2e):

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COMPOUND 26

phate. After cooling to 0 °C, a solution of phosphorus oxychloride (POCIs) in dry Compound (2d) (122 mg, 230 µmol) was dissolved in 400 µL dry trimethylphos-

- trimethylphosphate was added (400 µL stock solution (105 mg/mL), 276 µmol). The reaction mixture was stirred at 0 °C for 2h. ß
 - Subsequently a solution of tributylammonium pyrophosphate (273 mg, 576 µmol in added at 0 °C. The reaction was stirred at room temperature for 10 min. and then 1.2 mL dry DMF) and tributylamine (109 mg, 587 µmot in 600 µL dry DMF) was stopped by addition of 1.0 M triethylammonium hydrogencarbonate (1 mL). 2

Example 13: Preparation of Compound II

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COMPOUND

Removal of N-Boc protection group.

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- to pH = 1 using HCl and incubated at room temperature for 30 minutes. The mixture Following phosphorylation, 50 µl of the phosphorylation reaction mixture is adjusted is adjusted to pH 5.5 using equimolar NaOH and Na-acetate (pH 5.5) before purification on TLC.
- from the nucleotides derivatives using 100% methanol as running solution. Subse-From the crude mixture, 20 samples of 2 µl were spotted on kieselgel 60 F₂₅₄ TLC quently, the TLC plate is air-dried and the nucleotide-derivative identified by UV-(Merck). Organic solvents and non-phosphorylated nucleosides were separated Purification of nucleotide derivatives using thin-layer chromatography (TLC) 22
- shadowing. Kiesel containing the nucleotide-derivative was isolated and extracted wice using 10 mM Na-acetate (pH = 5.5) as solvent. Kiesetget was removed by ജ

centrifugation and the supernatant was dried *in vacuo*. The nucleotide derivative was resuspended in 50-100 µl H₂O to a final concentration of 1-3 mM. The concentration of each nucleotide derivative was evaluated by UV-absorption prior to use in polymerase extension reactions.

Examples 14 to 18: Preparation of the mononucleotide building block (III)

10 Building block III may be prepared according to the general scheme shown below:

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Example 14: Preparation of N-Boc-6-alanine propargyl amide(3a)

COMPOUND 2a

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Water was added (20 mL) and the product was extracted into EtOAc (3x30 mL). The combined EtOAc was dried (Na₂SO₄) and evaporated. The crude product was puri-N-Boc-β-alanine(1a) (1,05g, 5.5 mmol) and propargyl amine (0.90 g, 16.5 mmol) were dissolved in THF (10 mL). Diisopropyl-carbodiimide (DIC, 695 g, 5.5 mmol) fied by silica column chromatography eluting with EtOAc:Heptane gradient (2:3)was added and the reaction mixture was stirred for 16 h at room temperature. (3:2.5) (v/v). Product yield 0.925 g, 74 %.

'H NMR (CDC)₃) 8 6.69 (1H, bs, NH), 5,32 (1H, bs, NH), 4.04 (2H, bs), 3,41 (2H, dd), 2,45 (2H, t), 2.24 (1H, s), 1,44 (9H, s, 'Bu).

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Example 15: Preparation of compound (3b) 2

COMPOUND 3b

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A solution of 5-iodo-2'-deoxycytidine (176 mg, 0.5 mmol), N-Boc-β-alanine propargyl amide(14) and triethylamine (100 mg, 1.0 mmol) in dry DMF (5 mL) were stirred at room temperature. N2 was passed through the solution for 20 min.

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Fetrakis(triphenylphosphine)palladium(0) (66.5 mg, 0.057 mmol) and còpper(I) lodide (20.7 mg, 0.108 mmol) were added and the reaction mixture was stirred at room temperature for 5 d

ride (234 mg, 1.5 mmol) in anhydrous DMF (1 mL) was added and the resulting mix-Imidazole (112 mg, 1.6 mmol)was added. A solution of tert-butyldimethylsilyl chloture was stirred for 16 h at room temperature.

The reaction mixture was evaporated and EtOAc (25 mL) was added. The resulting mixture was filtrated and the solvent removed by vacuum evaporation.

The crude product was purified by silica column chromatography eluting with

DCM:MeOH (92.5-7.5) (v/v). Product yield 84 mg, 25%. 우

(9H, s, 'Bu), 0.95 (9H, s, 'Bu), 0.92 (9H, s, 'Bu), 0.17 (3H, s, CH₃), 0.15 (3H, s, CH₃), H NMR (CDC13) 8 8.13 (H, s), 6.21 (1H, dd, 1'-H), 4.66 (1H, m), 4,16 (2H, s, CH2), 4,04-3.85 (4H, m), 3.35-3.31 (2H, m), 2,43-2.36 (2H, m), 2.12-1.99 (1H, m), 1.44

0.13 (3H, s, CH₃), 0.12 (3H, s, CH₃). 15

Example 16: Preparation of compound (3c)

COMPOUND 3c

A solution of compound(3b) (84 mg, 0.12 mmol) and tetrabutylammonium fluoride trihydrate (TBAF) (155 mg, 0.45 mmol) in 2 mL dry THF was stirred at room tem-

perature for 4 days.

25

The reaction mixture was evaporated and purified by silica column chromatography sluting with DCM:MeOH gradient (9:1)-(8:2) (v/v). Product yield 27 mg, 48%.

3.95 (1H, q), 3.83 (1H, dd), 3.72 (1H, dd), 3,36-3.30 (3H, m), 2.42-2.36 (3H, m), 2.13 H NMR (CDCl₃) 8 8.32 (1H, s), 6.20 (1H, dd, 1'-H), 4.35 (1H, dt), 4,15 (2H, s, CH₂), (1H, dt), 1.40 (9H, s, 'Bu). ဗ္က

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Example 17: Preparation of compound (3d)

COMPOUND 3d

Compound (3c) (27 mg, 60 µmol) was dissolved in 100 µL dry trimethylphosphate.

phosphate was added (100 μL stock solution (110 mg/mL), 72 μmol). The reaction After cooling to 0 °C, a solution of phosphorus oxychloride (POCls) in dry trimethylmixture was stirred at 0 °C for 2h.

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Subsequently a solution of tributylammonium pyrophosphate (71 mg, 150 µmol ln added at 0 °C. The reaction was stirred at room temperature for 3 min. and then 300 µL dry DMF) and tributylamine (28.3 mg, 153 µmol in 150 µL dry DMF) was stopped by addition of 1.0 M triethylammonium hydrogencarbonate (1 mL).

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Example 18: Preparation of compound III

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Removal of N-Boc protection group.

to pH = 1 using HCl and incubated at room temperature for 30 minutes. The mixture Following phosphorylation, 50 µl of the phosphorylation reaction mixture is adjusted 22

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is adjusted to pH 5.5 using equimolar NaOH and Na-acetate (pH 5.5) before purification on TLC.

Purification of nucleotide derivatives using thin-layer chromatography (TLC)

- from the nucleotides derivatives using 100% methanol as running solution. Subseshadowing. Kiesel containing the nucleotide-derivative was isolated and extracted From the crude mixture, 20 samples of 2 µl were spotted on kieselgel 60 F₂₅₄ TLC quently, the TLC plate is air-dried and the nucleotide-derivative identified by UV-(Merck). Organic solvents and non-phosphorylated nucleosides were separated 'n
- tration of each nucleotide derivative was evaluated by UV-absorption prior to use in was resuspended in 50-100 μ l H $_2$ O to a final concentration of 1-3 mM. The concencentrifugation and the supernatant was dried in vacuo. The nucleotide derivative twice using 10 mM Na-acetate (pH = 5.5) as solvent. Kieselgel was removed by polymerase extension reactions. 9

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Examples 19 to 22: Preparation of the mononucleotide building block (IV)

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Building block IV may be prepared according to the general scheme shown below:

Example 19: Preparation of N-Acetyl-B-alanine(4a) S

COMPOUND 4a

To a solution of β-alanine (2,25 g, 25 mmol) in aq. NaHCO₃ (15 mL) was added acestirred at room temperature for 3 h. Acetic anhydride (2.55 g, 25 mmol) was added ionitrile (15 mL) and acetic anhydride (2.55 g, 25 mmol). The reaction mixture was and after 2 h and pH was adjusted to 4-5 by addition of NaH2PO4. 5

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The product was extracted into EtOAc (3 x 50 mL), dried (Na₂SO₄), and evaporated to dryness under vacuum to afford 1.96 g (60%)

Example 20: Preparation of N-Acetyl-B-alanine propargyl ester(4b).

COMPOUND 4b

dimethylaminopyridin (5 mg). The reaction mixture was stirred at room temperature To a solution of N-Acetyl-B-alanine(4a) in THF (20 mL) was added propargyl alcohol (840 mg, 15 mmol), 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride (EDC) (1.035 g.5.39 mmol), triethylamine (540 mg, 5.4 mmol) and 4-9

The reaction mixture was poured into EtOAc (100 mL), washed with NaH₂PO₄ (50% sat. aq, 2x50 mL) followed by NaHCO₃ (50% sat. aq, 50 mL). After drying (Na₂SO₄₎, EtOAc was removed under reduced pressure to leave a colourless oil that solidified on standing. Product yield 536 mg, 59%. 15

Example 21: Preparation of compound (4c)

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COMPOUND 4c

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A solution of 5-iodo-2'-deoxycytidin (200 mg, 0.56 mmol), triethylamine (100 mg, 1 mmol) and compound (4b) (190 mg, 1.13 mmol) in anhydrous DMF (7mL) was stirred at room temperature. N2 was passed through the solution for 20 min.

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Tetrakis(triphenylphosphine)palladium(0) (70mg, 0.06 mmol) and copper(1) lodide (22 mg, 0.12 mmol) were added and the reaction mixture was stirred at room temperature for 4 d.

The reaction mixture was evaporated and purified by silica column chromatography eluting with DCM:MeOH gradient (9:1)-(8:2) (v/v). Product yield 141 mg, 63%.

14 NMR (CD₃OD) & 8.41 (14, s), 6.20 (14, dd, 1'-H), 4.97 (2H, s), 4.38 (1H, dl), 3.97 (1H, q), 3.85 (1H, dd), 3.75 (1H, dd), 3.46 (2H, t), 2.61 (2H, t), 2.39 (1H, m), 2.18 (1H, m).

10 Example 22: Preparation of compound IV:

15 COMPOUND IV

Compound (4c) (140 mg, 355 µmol) was dissolved in 600 µL dry trimethylphosphate. After cooling to 0 °C, a solution of phosphorus oxychloride (POCIs) in dry trimethylphosphate was added (600 µL stock solution (108 mg/mL), 420 µmol). The reaction mixture was stirred at 0 °C for 2h.

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Subsequently a solution of tributylammonium pyrophosphate (422 mg, 890 µmol in 1.8 mL dry DMF) and tributylamine (168 mg, 900 µmol in 0.9 mL dry DMF) was added at 0 °C. The reaction was stirred at room temperature for 3 min. and then stopped by addition of 1.0 M triethylammonium hydrogencarbonate (1 mL).

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From the crude mixture, 20 samples of 2 µl were spotted on kieselgel 80 F₂₃₄ TLC (Merck). Organic solvents and non-phosphorylated nucleosides were separated from the nucleotides derivatives using 100% methanol as running solution. Subsequently, the TLC plate is air-dried and the nucleotide-derivative identified by UV-shadowing. Kiesel containing the nucleotide-derivative was isolated and extracted twice using 10 mM Na-acetate (pH = 5.5) as solvent. Kieselgel was removed by

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centrifugation and the supernatant was dried in vacuo. The nucleotide derivative was resuspended in 50-100 µl H2O to a final concentration of 1-3 mM. The concentration of each nucleotide derivative was evaluated by UV-absorption prior to use in polymerase extension reactions.

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Examples 23 to 27: Preparation of the mononucleotide building block (V)

Building block V may be prepared according to the general scheme shown below:

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Example 23: Preparation of 2-Aminoxy-acetic acid Ethyl ester (5a)

COMPOUND 5a

Acetyl chloride (5 mL) was added to abs. ethanole (50 mL) and the solution was

mmol) was added and the reaction mixture was stirred for 16 h at room temperature. product was extracted into diethyl ether (5x20 mL), dried (Na₂SO₄), and evaporated cooled to room temperature. 2-Aminoxy-acetic acid, hydrochloride (2:1) (1.10 g, 10 The reaction mixture was evaporated, K₂CO₃ aq. (2M) (10 mL) was added and the coold to afford 1.007 g, 84%.

¹H NMR (CDCl₃) 8 4.24 (2H, s), 4.22 (2H, q), 1.30 (3H, t). 9

Example 24: Preparation of Pent-4-ynoylaminooxy-acetic acid Ethyl ester (Sb)

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COMPOUND 5b

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Pentynoic acid (441 mg, 4.5 mmol) in 15 mL EtOAc were added dicyclohexylcarbodiimide (928 mg, 4.5 mmol) and the resulting mixture was stirred at room tem-To a solution of 2-Aminoxy-acetic acid ethyl ester (573 mg, 4.8 mmol) and 4perature for 16 h. The reaction mixture was filtered, and the filtrate was washed with EtOAc (2x5 mL). The combined EtOAc was washed with aq NaH2PO, and aq NaHCO3, dried (Na₂SO₄), and evaporated to afford 950 mg of crude product. 22

The crude product was purified by silica column chromatography eluting with EtOAc:Heptane gradient (1:3)-(1:1) (v/v). Product yield 700 mg, 78%

14 NMR (CDCl₃) 8 4.41 (2H, s), 4.18 (2H, q), 2.77 (1H, 1), 2.34 (2H, dt), 2.17 (2H, bt), 1.40 (3H, t). ဓ

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Example 25: Preparation of compound 5c.

COMPOUND 5c

A sotution of 7-Deaza-7-iodo-2'-deoxyadenosine (125 mg, 0.33 mmol), (prepared as described by Seela, F.; Synthesis 1996, 726-730), triethylamine (67 mg, 0.66 mmol) and compound(5b) (305 mg, 1.53 mmol) in anhydrous DMF (7mL) was stirred at room temperature. N₂ was passed through the solution for 20 min.

Tetrakis(triphenylphosphine)palladium(0) (75mg, 0.065 mmol) and copper(I) iodide (24 mg, 0.33 mmol) were added and the reaction mixture was stirred at room tem-

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perature for 16 h.

The reaction mixture was evaporated and purified by silica column chromatography eluting with DCM:MeOH (9:1) (vVy.) Product yield 129 mg, 86%.

H NMR (d° DMSO) § 11.6 (1H, s), 8.09 (1H, s), 7.63 (1H, s), 6.47 (1H, dd), 5.26 (1H, d), 5.08 (1H, t), 4.42 (2H, s), 4.32 (1H, m), 4.08 (2H, q), 3.81 (1H, m), 3.54 (2H, m), 2.66 (1H, t), 2.46 (1H, m), 2.30 (2H, t), 2.15 (2H, dd), 1.15 (3H, t).

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Example 26: Preparation of compound 5d:

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COMPOUND 5d

Compound (5c) (117 mg, 260 µmol) was dissolved in 500 µL dry trimethylphosphate. After cooling to 0 °C, a solution of phosphorus oxychloride (POCl₃) in dry

5 trimethylphosphate was added (400 µL stock solution (120 mg/mL), 310 µmol). The reaction mixture was stirred at 0 °C for 2h.
Subsequently a solution of tributylammoniumpyrophosphate (200 mg, 420 µmol in

1.00 mL dry DMF) and tributylamine (123.6 mg, 670 µmol in 500 µL dry DMF) was

added at 0 °C. The reaction was stirred at room temperature for 3 min. and then

10 stopped by addition of 1 mL 1.0 M triethylammonlumhydrogencarbonate.

Example 27: Preparation of compound V:

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COMPOUND V

The reaction mixture of compound (5d) (2.0 mL) was diluted with water (6.0 mL) and adjusted to pH 13 using NaOH (2M, aq). After incubation at 5 °C for 64 h, the reaction mixture was extracted with EtOAc (5x5 mL), adjusted to pH 7.0 using HCl (2M, aq), evaporated and diluted with triethylammonium acetate buffer (500 µL, 0.1 M ao).

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The crude product of triphosphate was purified by HPLC on a Waters Xterra MS C₁₈ Column, using the following buffer system: (A) aqueous triethylammonium acetate (0.1 M, pH 7) and (B) acetonitrile:water (80:20) containing triethylammonium acetate (0.1 M). The gradient time table contains 8 entries which are:

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Retention times of compound V and compound 5d were 4.82 min and 7.29 min respectively, measured by monitoring UV absorbance at 260 nm. The fractions containing pure product were pooled and lyophilized two times from water (3 mL).

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Examples 28 to 30: Preparation of the mononucleotide building block (VI)

Example 28: Preparation of Pent-4-ynoic acid 4-oxo-4H-

benzo[d][1,2,3]triazin-3-yl ester (6a) \$

2.04 mmol) in THF (2 mL) was added. 3-Hydroxy-1,2,3-benzotriazin-4(3H)-one (333 off. The filtrate was concentrated in vacuo and crystallized from hexane (4 mL), The Pentynoic acid (200 mg, 2.04 mmol) was dissolved in THF (4 mL). The solution was mg, 2.04 mmol) was added after 5 minutes. The reaction mixture was stirred 1h at -1,2,3-benzotriazin-4(3H)-one (eluent: ethyl acetate). Precipitated salts were filtered 10°C and then 2h at room temperature. TLC indicated full conversion of 3-hydroxycrystals were filtered off and dried. Yield: 450 mg, 93%. R_F = 0.8 (ethyl acetate). cooled in a brine-loawater bath. A solution of dicyclohexylcarbodiimide (421 mg,

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Example 29: Preparation of 2-Pent-4-ynoylamino-succinic acid 1-tert-butyl ester 4-isopropyl ester (6b)

was stirred overnight. Dichloromethane (10 mL) was added. The organic phase was L-Aspartic acid a, β-di-tert-butyl ester hydrochloride (Novabiochem 04-12-5066, 200 mg, 0.71 mmol) was dissolved in THF (5 mL). The activated ester 6a (173 mg, 0.71 mmol) and diisopropylethylamine (0.15 mL, 0.86 mmol) were added. The mixture

washed with citric acid (2 x 10 mL), saturated NaHCO₃ (aq, 10 mL), brine (10 mL), dried (Na₂SO₄) and concentrated to a syrup. An NMR spectrum indicated the syrup was pure enough for further synthesis. 1H-NMR (CDCls): 5 6.6 (1H, NH), 4.6 (1H, CH), 2.8 (2H, CH₂), 2.4 (4H, 2 x CH₂), 1.9 (1H, CH), 1.2 (18H, 6 x CH₃). 6

hydroxymethyl) tetrahydrofuran-2-yl) 2,4-dioxo-1,2,3,4-tetrahydro-pyrimidin-Example 30: Preparation of 2-(5-[1-(4-Hydroxy-5-(0-triphosphate-5-yl]-pent-4-ynoylamino}-succinic acid di-tert-butyl ester (VI)

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solution was degassed and kept under an atmosphere of argon. The catalyst The nucleotide (20 mg, 0.022 mmot) was dissolved in water-ethanol (1:1, 2 mL). The 2

Pd(PPh₂(m-C₆H₅SO₂Na⁺))₄ (20 mg, 0.016 mmol) prepared in accordance with A.L. Casalnuovo et al. J. Am. Chem. Soc. 1990, 112, 4324-4330, triethylamine (0.02 mL, 0.1 mmol) and the alkyne (Compound 6b) (20 mg, 0.061 mmol) were added. Few crystals of Cul were added. The reaction mixture was stirred for 6 h. The triethylammonium salt of LH8037 was achieved after purification by RP-HPLC (eluent: 100mM triethylammonium acetate → 20% acetonitrile in 100mM triethylammonium acetate). 'H-NMR (D₂O): δ 8.1 (1H, CH), 6.2 (1H, CH), 4.8 (1H, CH), 4.6 (1H, CH), 4.1 (3H, CH, CH₂), 2.8 (2H, CH₂), 2.7 (2H, CH₂), 2.5 (2H, CH₂), 2.3 (2H, CH₂), 1.4 (18H, 6 x CH₃).

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Immediately prior to incorporation or after incorporation, the protective di-tert-butyl ester groups may be cleaved to obtain the corresponding free carboxylic acid.

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Examples 31 to 32: Preparation of the mononucleotide building block (VII)

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Example 31: Preparation of 2-15-14-Amino-1-(4-hydroxy-5-hydroxymethyl-tetrahydrofuran-2-yl)-2-oxo-1,2-dihydro-pyrimidin-5-yl)-pent-4-ynoylamino-succinic acid di-tert-butyl ester (7a)

Compound (7a) (30 mg, 19%) was obtained from compound (6b) (140 mg, 0.43 mmol) and 5-iodo-2-deoxycytidine (100 mg, 0.28 mmol) using the procedure described for the synthesis of compound VI. ¹H-NIMR (MeOD-D₃): 6 9.3 (1H, CH), 6.2 (1H, CH), 4.8 (1H, CH), 4.6 (1H, CH), 4.4 (1H, CH), 4.0 (1H, CH), 3.8 (2H, CH₂), 2.7 (1H, CH₂), 2.2 (1H, CH₂), 1.4 (18H, 6 x CH₃).

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Example 32: Preparation of 2-(5-14-Amino-1-(4-hydroxy-5-(0-triphosphate-hydroxymethyl)-tetrahydro-furan-2-yl)-2-oxo-1,2-dihydro-pyrimidin-5-yll-pent-4-noylamino)-succinic acid di-tert-butyl ester (Compound VII)

Phosphoroxy chloride (6.0 µl, 0.059 mmol) was added to a cooled solution (0°C) of 7a (30 mg, 0.054 mmol) in trimethyl phosphate (1 mL). The mixture was stirred for 1h. A solution of bis-n-tributylammonium pyrophosphate (77 mg, 0.16 mmol) in DMF (1 mL) and tributylamine (40 µl, 0.16 mmol) were added. Water (2 mL) was added, pH of the solution was measured to be neutral. The solution was stirred at room temperature for 3 h and at 5 °C overnight. A small amount of compound VII (few mg) was obtained after purification by RP-HPLC (eluent: 100mM triethylammonium acetate → 20% acetonitrile in 100mM triethylammonium acetate). 7a (18 mg) was regained.

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Immediately prior to or subsequent to incorporation the protective di-tert-butyl ester groups may be cleaved to obtain the corresponding free carboxylic acid.

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Examples 33 and 34: Preparation of the mononucleotide building block

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Example 33: Preparation of 2-Pent-4-ynoylamino-6-(2,2,2-trifluoro-acetylamino-hexanoic acid, (8a)

Compound 6a (250 mg, 1.0 mmol) was added to a solution of *N*-ɛ-trifloroacety,L-lysine (Novablochem, 04-12-5245) (250 mg, 1.0 mmol) in DMF (3 mL). Ethyldiisopropylamine (0.2 mL, 1.2 mmol) was added. The solution was stirred at room temperature overnight and worked-up by RP-HPLC (eluent: water → methanol). Yield: 50 mg, 15% 'H-NMR (0₂O): 6 4.4 (1H, CH), 3.4 (2H, CH₂), 2.5 (4H, 2 x CH₂), 2.3 (1H, CH), 1.9 (1H, CH₂), 1.8 (1H, CH₂) 1.6 (2H, CH₂), 1.5 (2H, CH₂).

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Example 34: Preparation of 2-(5-(1-(4-Hydroxy-5-(O-triphosphate-hydroxymethyl)-tetrahydrofuran-2-yl)-2,4-dioxo-1,2,3,4-tetrahydro-pyrimidin-5-yl)-pent-4-ynoylamino}-6-(2,2,2-trifluoro-acetylamino}-hexanoic acid_(Compound VIII)

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The triethylammonium salt of compound VIII (11 mg) was obtained from compound 8a (50 mg, 0.15 mmol) and 5-iodo-2-deoxyuracil (50 mg, 0.06 mmol) using the procedure described for the synthesis of compound VI.

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Examples 35 to 39: Preparation of the mononucleotide building block (IX)

Example 35: Preparation of di-Boc-Lysin-propargyl amide (compound9a)

C₁₈H₃₃N₃O₅ Mw 383.48

Boc-Lys-(Boc)-OSu (Novabiochem 04-12-0017, 0.887 g, 2 mmol) was dissolved in THF (10 ml). Propargylamine (0.412 ml, 6 mmol) was added and the solution stirred for 2 h. TLC (ethylacetate:heptan 1:1) showed only one product. Dichloromethane (20 ml) was added and the mixture was washed successively with citric acid (1M, 10 ml) and saturated sodium hydrogen carbonate (10 ml). The organic phase was dried with magnesium sulphate filtered and evaporated to give compound 9a (0.730 g) as a colourless syrup.

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'H-NMR: ϑ 6.55 (1H, NH), 5.15 (1H, NH), 4.6 (1H, $\underline{\text{CH}}$ -NH), 4.05 (2H, CH-C- $\underline{\text{CH}}$ -N), 3.75 (1H, NH), 3.1 (2H, $\underline{\text{CH}}$ -NH) 2.25 (1H, $\underline{\text{CH}}$ -C-CH₂), 1.9-1.3 (6H, 3 × CH₂), 1.4 (18H, 6 × CH₃).

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20 Example 36: Preparation of 5-lodo-3'-O-acetyl-5'-O-TBDMS-2'-deoxyuridine (compound 9b) C₁₇H₂₂IN₂O₈Si Mw 510.40

ure. The reaction mixture was evaporated and dissolved in dichloromethane (20 ml) TBDMSCI (t-butyl-dimethyl-chloride, 1,12 g, 7.41 mmol) in dichloromethane (5.0 ml) dichloromethane (2 x 20 ml). The combined organic phases were washed with saturated sodium bicarbonate (20 ml), dried with sodium sulphate and evaporated (5.85 and citric acid (2M, 20 ml) was added. The aqueous phase was back extracted with solved in pyridine (40 ml) and cooled to 0 °C. Acetic anhydride (4.0 ml, 42.3 mmol) was run in over 20 minutes. Stirring was continued at room temperature for 18 h, was added over 30 minutes and stirring was continued for 18 h at room tempera-5-lodo-2'-deoxyuridine (Sigma I-7125, 2.50 g, 7.06 mmol) and imidazol (0.961 g, g). Recrystallisation form ethylacetate/EtOH gave 9b (2.54, g) pure for synthesis TLC (Ethyl acetate). Further recrystallisation furnished an analytical pure sample and the mixture was evaporated. The crude mono silylated nucleoside was dis-14.12 mmol) was dissolved in DMF (10 ml). Cooled to 0 °C and a solution of mp.172.4-173.1 °C.

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Example 37: Preparation of 5-lodo-3'-O-acetyl-2'-deoxyuridine (compound 9c) C₁₁H₁₃IN₂O₆ Mw 396.14

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5-lodo-3'-O-acetyl-5'-O-TBDMS-2'-deoxyuridine (compound 9b) (2.54 g, 4.98 mmol) reduced to approximately 10 ml in vaccuo. Crystals were collected and dried in vac-10.1 mmol) was added and stirred for 18 h at room temperature. The reaction mixture was added water (25 ml) stirred for 1 h. Ion exchange resin IR-120 H* (26 ml) as dissolved in THF (25 ml), tetra butyl ammonium fluoride trihydrat (TBAF, 3.2 g. was then added and stirring was continued for 1 h. The solution was filtered and

Example 38: Preparation of 5-lodo-5'-triphosphate-2'-deoxyuridine, triethylammonium salt (compound 9d) C₉H₁₄IN₂O₁₄P₃ + n·N(CH₂CH₃)₃ Mw 897.61 for n =3. 5

was continued for 10 minutes and the intermediate was oxidized by adding an lodine DMF (9.81 ml, 0.5 M, 4.91 mmol) and tri-n-butylamine (3.12 ml, 13.1 mmol). Stirring 5-lodo-3'-O-acetyl-2'-deoxyuridine (compound 9c) (2.54 g. 4.98 mmol) as dissolved benzodioxaphosphorin-4-one in dioxane (3.60 ml, 1 M, 3.60 mmol) was added unsolution (90 ml, 1% w/v in pyridine/water (98/2, v/v)) until permanent iodine colour. followed by simultaneous addition of bis(tri-n-butylammonium) pyrophosphate in der stirring. The reaction mixture was stirred for 10 minutes at room temperature in pyridine (3.2 ml) and dioxane (10 ml). A solution of 2-chloro-4H-1,3,2-15 8

he reaction mixture was left for 15 minutes and then decolourized with sodium thiosulfate (5% aqueous solution, w/v). The reaction mixture was evaporated to yellow ammonia (100 ml, 25%) was added. This mixture was stirred for 1.5 hour at room emperature and then evaporated to an oil of the crude triphosphate product. The oil. The oil was stirred in water (20 ml) for 30 minutes and concentrated aqueous 22

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crude material was purified using a DEAE Sephadex A25 column (approximately 100 ml) eluted with a linear gradient of triethyl- ammonium hydrogencarbonate [TEAB] from 0.05 M to 1.0 M (pH approximately 7.0 – 7.5); flow 8 ml/fraction/15 minutes. The positive fractions were identified by RP18 HPLC eluting with a gradient from 10 mM TEAA (triethylammonium acetate) in water to 10 mM TEAA 20% water in acetonitrile. The appropriate fractions were pooled and evaporated. Yield approximately 1042 mg.

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Example 39: Preparation of 5-(Lysin-proparayl amide)-5'-triphosphate-2'-deoxycytidine, triethylammonium salt (compound IX) C₁₈H₃₀N₅O₁₅P₃ + n-N(CH₂CH₃)₃ Mw 952.95 for n = 3

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5-lodo-3-Co-acetyl-5-triphosphate-2'-deoxyuridine, triethylammonium salt (compound 9d) (0.0087 g, 9.7 µmol) was dissolved in water (100 µl). Air was replaced carefully with argon. Di-Boc-Lysin-propargyl amide (compound 9a) (18.6 mg, 48.5 µmol) dissolved in dioxane (100 µl), triethylamine (2.7 µl, 19.4 µl), Pd((PPh₂)(m-C₆H₄SO₃Na²)*(H₂O))₄ (compound 9d) (5 mg, 4.4 µmol) and copper (l) iodide (0.4 µl, 2.1 µmol) were added in the given order. The reaction mixture was stirred for 18 h at room temperature in an inert atmosphere then evaporated. The crude material was treated with aqueous hydrochloric acid (0.2 M, 1 ml) for 15 minutes at 30 °C. (compound IX) was obtained by HPLC C₁₆ 10 mM TEAA (triethylammonium acetate) in water to 10 mM TEAA 20% water in acetonitrile. Appropriate fractions were desalted using gelfiltration (pharmacia G-10, 0.7 ml).

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Examples 40 to 45: Preparation of the mononucleotide building block (X)

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Example 40: Preparation of Boc-Lys-(Boc)-OH (compound 10a) C₁₆H_{3g}N₂O₆ Mw 346.42

Lysine (Novabiochem 04-10-0024; 3.65 g, 20 mmol) was dissolved in sodium hydroxide (2 M, 40 ml), added dioxane (80 ml) and di-tert-butyl dicarbonate (8.73 g, 40 mmol) in the given order. The mixture was stirred for 1.75 h at 60 °C. Water (50 ml) was added and the solution was washed with dichloromethane (4 x 25 ml). The aqueous phase was cooled to 0 °C with ice then acidified with 2 M HCl (pH = 3) and extracted with dichloromethane (4 x 25 ml). The organic phase was dried with mag-

10 nesium sulphate. Evaporation fumished (compound 10a) 6.8 g as a colour less oil. 'H-NMR: 3 9.5 (1H, COOH), 5.3 (1H, CH), 4.7 (1H, NH), 4.3 (1H, NH), 3.1 (2H, CH₂-NH), 1.8 (2H, CH₂-CH), 1.5(6H, 3xCH₂), 1.45 (18H, 6 x CH₃).

15 Example 41: Preparation of di-Boc-Lysin-propargyl ester (compound 10b) C₁₉H₃₂N₂O₆ Mw 384.47

20 Boc-Lys-(Boc)-OH (compound 10a) (3.46 g, 10 mmol) was dissolved in THF (25 ml). At 0 °C a solution of dicyclohexylcarbodimide (2.02 g, 10 mmol) in THF (25 ml) and triethylamine (1.39 ml) were added in the given order. The mixture was allowed to warm up to room temperature and stirred for 18 h. The resulting suspension was filtered and evaporated. The oil 5.45 g was pre-purified by column chromatography Heptan: Ethylacetate 3:1.

Pure 10b was achieved by HPLC- C₁₆ 10% MeOH: 90% H₂O → 100% MeOH

'H-NMR: 35.1 (1H, NH), 4.75 (2H, CH-C-CH2-O), 4.6 (1H, NH), 4.35 (1H, CH-NH), 3.1 (2H, CH₂-NH) 2.5 (1H, CH-C-CH₂), 1.9-1.4 (6H, 3 x CH₂), 1.5 (18H, 6 x CH₃). Example 42: Preparation of 5-lodo-3',5'-di-O-TBDMS-2'deoxycytidine (com-

pound 10c) C21H40IN3O4Si2 Mw 581.64 2

The combined organic phases were washed with saturated sodium bicarbonate (15 5-lodo-2-deoxy-Cytidine (Sigma I -7000, 0.353 g, 1 mmol) was dissolved in DMF (4 ml), added t-Butyl-dimethyl silyl chloride (TBDMS-Cl, 0.332 g, 2.2 mmol) and Imidazol (0.204 g, 3 mmol). The solution was stirred for 15 h at 50 °C followed by evapomixture. The aqueous phase was back extracted with dichloromethane (2 \times 10 ml). ml), dried with sodium sulphate and evaporated. Compound 10 c (0.405 g) was obration. Dichloromethane (25 ml) and citric acid (2M, 10 ml) was added to the dry tained by recrystallisation from EtOH/Ethylacetate.

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'H-NMR: 38.1 (1H, H-6), 6.25 (1H, H-1'), 4.35 (1H, H-4'), 4.0 (1H, H-4'), 3.9 (1H, H-5), 3.75 (1H, H-5'), 2.5 (1H, H-2'), 1.95 (1H, H-2'), 1.85 (2H, NH), 0.95 (9H, 3 x CH₃), 0.9 (9H, 3 × CH₃), 0.15 (6H, 2 × CH₃), 0.1 (6H, 2 × CH₃).

Preparation of 5-(di-Boc-Lysin-propargyl ester)-3',5'-di-O-TBDMS-2'deoxycytidine (compound 10d) C40H71IN5O10Siz Mw 838.19

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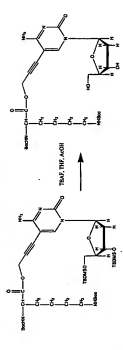
Compound 10c (0.116 g, 0.2 mmol) was dissolved in dichloromethane (10 ml). Air was replaced carefully with argon. Di-Boc-Lysin-propargyl ester (compound 10b) (0.232, 0.6 mmol), triethylamine (0.083 ml, 0.6 mmol), di-chloro-bis-S

g, 0.02 mmol) were added in the given order. The reaction mixture was stirred for 15 triphenylphosphine-palladium II (0.0074 g, 0.01 mmol) and copper (I) iodide (0.0038 h at room temperature in an inert atmosphere. The reaction mixture was evaporated re-dissolved in MeOH/H₂O 1:1 1 ml and purified using HPLC-C₁₈ 45% H₂O:55% MeCN → 100% MeCN. 9

H-NMR: θ ¹H-NMR: θ 8.2 (1H, H-6), 6.25 (1H, H-1'), 5.15 (1H, NH), 4.9 (2H, C- $\overline{\text{CH}}_{Z}$ CH₂), 1.85 (2H, NH), 1.5 (18H, 6 x CH₃), 0.95 (9H, 3 x CH₃), 0.9 (9H, 3 x CH₃), 0.15 3.75 (1H, H-5'), 2.5 (1H, H-2'), 3.1 (2H, CH2-NH), 1.95 (1H, H-2'), 1.9-1.4 (6H, 3 x O), 4.6 (1H, NH), 4.4 (1H, H-4'), 4.3 (1H, CH-NH), 4.0 (1H, H-4'), 3.9 (1H, H-5'), (6H, 2 x CH₃), 0.1 (6H, 2 x CH₃).

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Example 44: Preparation of 5-(di-Boc-Lysin-propargyl ester)-2'-deoxycytidine (compound 10e) C₂₈H₄₃IN₅O₁₀ Mw 609.67



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Compound 10d (0.0246 g, 0.029 mmol) was dissolved in THF (1 ml) and successively added acetic acid (0.0165 ml, 0.288 mmol) and tetra n-butyl ammonium fluodide tri-hydrate (0.0454 g, 0.144 mmol). The reaction mixture was stirred for 18 h at room temperature and afterwards evaporated. Re-dissolved in dichloromethane and purified on silica (1 x 18 cm). Dichloromethane/MeOH 8:2. Fractions which gave UV absorbance on TLC were pooled and evaporated giving (0.0128 g) as a colourless oil.

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Example 45: Preparation of 5-(Lysin-propargyl ester)-5'-triphosphate-2'-

10 deoxycytidine C₁₈H₃₀N₅O₁₅P₃ Mw 649.38

Compound 10e (0.0128 g, 0.021 mmol) was dissolved in trimethylphosphate (0.150 ml) and cooled to 0 °C. Phosphoroxychloride in trimethylphosphate (1M, 0.0246 ml) was added slowly in order not to raise the temperature. Stirring was continued for 2 h at 0 °C and the temperature was allowed to rise to ambient. Pyrophosphate in DMF (0.5 M, 0.1025 ml, 0.051 mmol) and tri-n-butyl amine in DMF (1M, 0.0122 ml, 0.051 mmol) were added stirrultaneous. Stirring was continued for 15 minutes at room temperature and TEAB(triethyl ammonium blcarbonate, 1M, pH = 7.3, 0.50ml) was added. Stirring was continued for 3 h then evaporated.

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Example 46: Preparation of compound X

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The crude material was treated with aqueous hydrochloric acid (0.2 M, 1 ml) for 15 minutes at 30 °C. Compound X was obtained by HPLC C₁₈ 10 mM TEAA (triethylamnonlum acetate) in water to 10 mM TEAA 20% water in acetonitrile. Appropriate fractions were desalted using gelfiltration (pharmacia G-10, 0.7 ml)

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Examples 47 to 51: Preparation of the mononucleotide building block (XI)

Example 47: Preparation of 3'-0-acetyl-5'-0-dimethoxytrityl-5-lodo-2'- deoxyuridine (compound 11a). C₃₂H_{JJ} IN₂0₈. Mw 698.51 g/mol. (Analogous to "Oligonucleotide Synthesis - a practical approach" (1984) Gait, M.J. (Ed.), IRL Press, Oxford.)

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5-lodo-2'-deoxyuridine (3.54 g, 10 mmol) was dried by coevaporation with pyridine (25 ml, 3 times). Pyridine (100 ml) was added and shortly evaporated to a reduced

DMT-CI (0.68 g, 2 mmol) was added and the reaction mixture was stirred for anothe and the reaction mixture was stirred at room temperature. After 20 hours, additlonal volume (80 ml). 4,4*-dimethoxytrityl chloride (DMT-Cl, 3.38 g, 10 mmol) was added 4 hour. Excess of DHT-CI was quenched with methanol (5 ml, stirred 10 minutes)

lions of dichloromethane were dried with anhydrous MgSO4, filtered and evaporated on silica (Merck Kieselgel 60, 230-400 mesh ASTM, art. 9385) eluting with a gradi-The crude oil was dissolved in dichloromethane (75 ml) and triturated with pentane odo-2'-deoxyuridine was obtained via column chromatography in dichloromethane and the reaction mixture evaporated to dryness. The oil was dissolved in dichloroadding pentane (250 ml) gave reddish foam after evaporation. Yield of crude 5'-0aqueous phase was back-extracted with dichloromethane and the combined frac-(250 ml). Re-trituration of the crude oil by dissolving in ethyl acetate (75 ml) and dimethoxytrityl-5-iodo-2'-deoxyuridine was, 5.84 g. Pure S'-O- dimethoxytrityl-Smethane (100 ml) and extracted with saturated aqueous NaHCO₃ (100 ml). The ent of methanol (0-5 % methanol in dichloromethane). Yield of purified 5'-0dimethoxytrltyl-5-iodo-2'-deoxyuridine was 4.26 g (6.5 mmol, 65%). S 5

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extracted with dichloromethane and the combined fractions of dichloromethane were 5'-0-Dimethoxytrityl-5-iodo-2'-deoxyuridine (6.0 g, 9.1 mmol) was dried by coevaporation with pyridine (10 ml, twice). Pyridine (50 ml) was added and acetic anhydride 5 ml) and dimethylaminopyridine (DMAP, catalytic amount) were added. The reaction mixture was stirred overnight at room temperature. Excess of acetic anhydride dimethoxytrityl-5-lodo-2'-deoxyuridine was. obtained via column chromatography in evaporated to dryness The oil was dissolved in dichloromethane (150 mt) and exracted with aqueous saturated NaHC03 (50 ml). The aqueous phase was backdried with anhydrous MgSOs, filtered and evaporated. Purified 3'-O-acetyl-5'-Owas quenched with methanol (10 ml, stirred 15 min.) and the reaction mixture ឧ 22

desired purified 3'-0-acetyl-5'-0-dimethoxytrityl-5-iodo-2'-deoxyuridine (4.18 g, 6.0 methane/pentane (80/20, v/v) eluting with a gradient of methanol (2-6') gave the. methane). The yield was 5.75 g (8.2 mmol). Rechromatography in dichlorommol, 60%) ဗ္က

dichloromethane/methanol (98/2, v/v) on silica (Merck Kieselgel 60, 230-400 mesh

ASTM, art.9385) eluting with a gradient of methanol (2-6 % methanol in dichloro-

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11b). C₅H₄F₃NO. Mw. 151.09 q/mol. (Reference: Cruickshank et al. (1988) Example 48: Preparation of N-trifluoroacetyl-3-amidopropyne (compound Tetrahedron Lett. 29, 5221-5224)

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Propargylamine (7.0 ml, 5.88 g, 0.11 mol) was dissolved in 100 ml ice-cold methanol and ethyl trifluoroacetate (18 ml, 19.2 g, 0.135 mol) was added slowly under stirring

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propargylamine (as observed by disappearance of the positive colour-reaction in the on ice. The ice bath was removed and the reaction mixture was allowed to warm up to room temperature and stirring v as continued over night. After 24 h, TLC analysis The aqueous phase was back-extracted with dichloromethane (25 ml) and the comchloromethane phases were dried with magnesium sulfate, filtered and evaporated chloromethane (100 ml) and extracted with aqueous sodium hydrogen carbonate. bined dichloromethane phases were extracted with water (100 ml). The aqueous ninhydrin test, 110 °C). The reaction mixture was evaporated, re-dissolved in diphase was back-extracted with dichloromethane (25 mi) and the combined di-(Silica, dichloromethane/methanol, 9/1, v/v) shoved complete conversion of

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to yellow oil. The oil was purified by distillation collecting the purified product at 38-

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39 °C/ 1mmHg. Yield 11.0 g (73 mmol, 66%)

acetvi-3-amido-propynyl)-2'-deoxyuridine (compound 11c), C35H35N309, Mw Example 49: Preparation of 3.-O-acetyl-5'-O-dimethoxytrityl-5-(N-trifluoro-625.67 g/mol

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rifluoroacetyl-3-amido-propynyl)-2'-deoxyuridine (4.2 g) as brownish oil. Rechromaaqueous EDTA (5% v/v, 300 ml) and once with aqueous sodium thiosulfate (5% v/v, rated. Column chromatography in dichloromethane/pentane (80/20, v/v) eluting with TLC analysis (Silica, CH2Cl2/MEOH, 95/5, v/v) and stopped after 24 hours when all a gradient of methanol (0-5%) gave the crude 3'-O-acetyl-5'-O-dimethoxytrityl-5-(N-300 ml). The aqueous phases were back-extracted with ethyl acetate and the com-3'-O-Acetyl-5'-O-dimethoxytrityl-5-iodo-2'-deoxyuridine (4.15 g, 6.0 mmol) was dissolved in ethyl acetate (240 ml) and N- trifluoroacetyl-3-aminopropyne (1.81 g, 12 mmol) were added in the given order. The reaction mixture was flushed with nitrogen, Stoppered and stirred at ambient temperature. The reaction was followed by bined fractions of ethyl acetat. were dried (anhydrous MgSO4), filtered and evapotography in ethylacetate/pentane (50/50 to 60/40, v/v) gave the desired purified starting material was consumed. The reaction mixture was extracted twice with palladium(11) chloride (0.091 g, 0.13 mmol) and copper(1) iodide (0.091 g, 0.48 mmol), triethylamine (3.09 g, 4.23 ml, 30.5 mmol), bis(triphenylphosphine)product (1.99 g, 3.2 mmol).

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Example 50: Preparation of 3'-O-acetyl-5-(N-trifluoroacetyl-3-amidopropynyl)-2'-deoxyuridin.. C16H16F3N307, Mw 419.31 g/mol.

drous MgSO₄), filtered and evaporated. The foam was dissolved in dichloromethane added slowly and the reaction mixture was stirred for 15 min at 0 °C. TLC analysis and the reaction mixture was poured into saturated aqueous NaHCO₃ (100 ml) and extracted twice with dichloromethane. The aqueous phase was back-extracted with dichloromethane and the combined fractions of dichloromethane were dried (anhyquenched by the addition of 2-propanol (10 ml), quenching of DMT was observed cooled to O °C. A solution of trichloroacetic acid in dichloromethane (3% w/v) was Silica, CH₂Cl₂/MeOH, 95/5 v/v) confirmed total detritylation and the reaction was by colour-change from orange to colourless. Stirring vas continued for 2 minuttes deoxyuridine (1.99 g, 2.8 mmol) was dissolved in dichloromethane (133 ml) and 3'-O-Acetyl-5'-O-dimethoxytrityl-5-(N-trifluoroacetyl-3-amidopropynyl)-2'-9 5 8

precipitate was redissolved and evaporated, first from methanol and then from chloafter rechromatography, eluting with a gradient in dichloromethane/methanol (98/2 (50 mt) and triturated with pentane (200 ml). The trituration was repeated and the amidopropynyl)-2'-deoxyuridine was obtained by silica gel column chromatography n dichloromethane/methanol (gradient: 95/5 to 89/11, v/v), The yield was. 0.37 g roform to give yellow foam. Purified 3'-O-acetyl-5-(N-trifluoroaoctyl-3-

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Example 51: Preparation of 5-(3-aminopropyl)-5'-triphosphate-2'-deoxyuridine, triethylammonium salt (compound XI), C₁₂H₁₈N₃O₁₄P_{3.+}n·N(CH₂CH₃)₃. Mw 824.78 g/mol for n = 3. (Ludwig, J. and Eckstein, F. (1989) J. Org. Chem. 54, 631-635).

solved in anhydrous pyridine (100 µl) and anhydrous dioxane (300 µl). A solution of for 10 minutes and the intermediate was oxidized by adding an iodine solution (3 ml, 1% w/v in pyridine/water (98/2, v/v)) until permanent iodine colour. The reaction mixflask (50 ml) with water and evaporated to yellow oil. The oil was stirred in water (10 oil of the crude triphosphate product. The crude material was purified using a DEAE mately 7.5 - 8.0); flow 8 ml/fraction/15 minutes. The positive fractions were identified 2-chloro-4H-1,3,2-benzodioxaphosphorin-4-one in dioxane (110 µl, 1 M, 0.11 mmol) was added under stirring and after 30 seconds precipitation of pyridinium hydrochloure was left for 15 minutes and then decolourized with sodium thiosulfate (4 drops, mmol) was dissolved in anhydrous pyridine (2 ml) and evaporated. The oil was dis-5% aqueous solution, w/v). The reaction mixture was transferred to a roundbottom This mixture was stirred for 1 hour at room temperature and then evaporated to an phate in DMF (300 µl, 0.5.M) and tri-n-butylamine (100 µl). Stirring was continued triethyl- ammonium hydrogencarbonate [TEAB] from 0.05 M to 1.0 M (pH approxiperature followed by simultaneous addition of bis(tri-n-butylammonium) pyrophosml) for 30 minutes and concentrated aqueous ammonia (20 ml, 32%) was added. 3'-O-Acetyl-5-(N-trifluoroacetyl-3-amidopropynyl)-2'-deoxyuridine (42.5 mg, 0.10 ride was observed. The reaction mixture was stirred for 10 minutes at room tem-Sephadex A25 column (approximately 100 ml) eluted with a linear gradient of

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CH₂), 1.4 (9H, 3 x CH₃).

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by RP18 HPLC eluting with a gradient from 10 mM TEAA (triethylammonlum acetate) in water to 10 mM TEAA 20% water in acetonitrile. The appropriate fractions were pooled and evaporated. Yield approximately 90 mg.

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Examples 52 to 54: Preparation of the mononucleotide building block (XII)

Example 52: Succinic acid mono-(3-tert-butoxycarbonylamino-propyl) ester [Compound 12a]

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Triethylamine (5.0 mL, 36 mmol) and di-tert-butyl dicarbonate (7.0 g, 32 mmol) were added to a solution of 3-aminopropanol (1.0 g, 26.6 mmol) in methanol (10 mL). The solution was stirred for 2 h. at room temperature. Methanol was evaporated off and the residue was dissolved in water (50 mL) and extracted with dichloromethane (50 mL). The organic phase was dried (Na₂SO₄) and concentrated in vacuo. The crude material was dissolved in dichloromethane (20 mL) and DMF (4 mL). Triethylamine (5.0 mL, 36 mmol) and succinic anhydride (3.0 g, 30 mmol) were added portion wise to the solution (exothermic reaction). The reaction mixture was stirred for 2 h, then concentrated and worked-up by RP-HPLC (eluent: water → methanol). Yield 6.0 g, 82%. 'H-NMR (CDCi₃): 6 4.2 (2H, CH₂), 3.2 (2H, CH₂), 2.7 (4H, 2 x CH₂), 1.8 (2H,

9-[4-(Isopropyl-dimethyl-silanyloxyl-5-(isopropyl-dimethyl-silanyloxymethyl)tetrahydro-furan-2-yll-9H-purin-6-ylamine (compound 12b)

Imidazole (2.0 g, 29.4 mmol) and tert-butyldimethylsilyl chloride (3.0 g, 19.9 mmol) were added to a solution of deoxyadenosine monohydrate (1.33, 4.94 mmol) in DMF to a solid in vacuo. Work-up by flash chromatography afforded crystalline compound 12b in a yield of 2.1 g, 94%. 'H-NMR (CDCI₃): 5 8.3 (1H, HC=), 8.1 (1H, HC=), 6.4 (1H, CH), 6.0 (2H, 2 x OH), 4.6 (1H, CH), 4.1 (1H, CH), 3.9 (1H, CH), 3.8 (1H, CH), (10 mL). The solution was stirred at 60°C overnight. The mixture was concentrated 2.6 (1H, CH₂), 2.4 (1H, CH₂), 0.9 (18H, 6 × CH₃), 0.0 (12H, 4 × CH₃).

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purin-6-yll-succinamic acid 3-tert-butoxycarbonylamino-propyl ester (com-Example 53: N-19-(4-Hydroxy-5-hydroxymethyl-tetrahydrofuran-2-vl)-9Hpound 12d

was added to an ice-water cooled solution of 12a (488 mg, 1.78 mmol) in THF (10 was washed with saturated NaHCO₃ (20 mL), dried (Na₂SO₄) and concentrated to a CH₂), 4.1 (1H, CH), 3.8 (1H, CH), 3.7 (2H, CH₂), 3.0 (4H, 2 × CH₂), 2.7 (3H, 2 × A solution of dicyclohexylcarbodilmide (366 mg, 1.78 mmol) in ethyl acetate (15 mL) ml.). Few crystals of 4-dimethylaminopyridine and 12b (850 mg, 1.78 mmol) were added. The reaction temperature was slowly raised to room temperature and the mixture was stirred overnight. Precipitated salts were filtered off. The organic phase solid. Approximately 20 mg of 12c was isolated after flash chromatography and 510 mg of starting material 12b was regained. 12c (20 mg) was dissolved in THF (2 mL). Tetrabutylammonium fluoride, trihydrate (100 mg) and acetic acid (0.2 mL) were added. The mixture was stirred for 1 day, then concentrated in vacuo and worked-up by column chromatography. Yield 10 mg. Compound 12c 1H-NMR (CDCI3): 5 8.4 (1H, HC=), 8.2 (1H, HC=), 6.4 (1H, CH), 5.8 (2H, 2 x OH), 4.6 (1H, CH), 4.2 (2H, CH₂), 2.4 (1H, CH₂), 1.8 (2H, CH₂), 1.4 (9H, 3 × CH₃), 0.9 (18H, 6 × CH₃), 0.0 (12H, 9 5

4 × CH₃). Selected NMR data for 12d: ¹H-NMR (MeOD-D₃): δ 4.6 (14, CH), 4.1 (2H, CH₂), 3.8 (1H, CH), 3.7 (1H, CH), 3.6 (2H, CH₂), 3.2 (2H, CH₂), 3.0 (2H, CH₂). 2.8 (3H, 2 × CH₃), 2.5 (1H, CH₂), 1.8 (2H, CH₃), 1.4 (9H, 3 × CH₃).

Example 54; N-19-(4-Hydroxy-5-hydroxymethyl-tetrahydrofuran-2-yl)-9H-purin-6-yl]-succinamic acid 3-tert-butoxycarbonylamino-propyl ester (compound XII)

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LH8075b (10 mg) was converted to the corresponding triphosphate LH8075c using the procedure described for the synthesis of compound VII. TLC indicated full conversion of compound 12d.

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Immediately prior to incorporation, the tert-butoxy group may be hydrolysed to release the free carboxylic acid. Alternatively, the tert-butoxy group may be cleaved after the formation of the templated molecule.

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Example 55: N-[9-(4-Hydroxy-5-(O-triphosphate-hydroxymethyl)-

20 tetrahydrofuran-2-yl)-9H-purin-6-yl]-succinamic acid (Compound XIII)

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dATP (5 µmol) was suspended in DMF (4 x 1 mL) and concentrated to a solld *in vacuo* four times. The solid was suspended in DMF (1 mL). Succinic anhydride (5 mg, 0.05 mmol) was added at -20°C. The mixture was stirred for 3h, and then concentrated to solid and purified by RP-HPLC (eluent: 0.1 % HCOOH in water → 10% methanol, 0.1% HCOOH in water). The purified material was dissolved in aqueous ammonia (25%, 1 mL) and stirred for 3h. The mixture was concentrated *in vacuo* and worked-up by RP-HPLC (eluent: 0.1 % HCOOH in water → 10% methanol, 0.1% HCOOH in water). Comparison with starting material indicated that the product

Examples 56 and 57: Preparation of the mononucleotide building block (XIV)

sluted 40s later off the column than the starting material.

15 Example 56: Benzyloxy-ethynyl-diisopropyl-silane (Compound 14a)

A solution of benzyl alcohol (0.1 mL, 1.0 mmol) in THF (0.5 mL) was added dropwise to a cooled (-78°C) solution of disopropylethylamine (1 mL), dichlorodiisopropyleilane (0.3 mL, 1.62 mmol) in THF (4 mL). The solution was stirred for 3h (-78 \rightarrow -20°C). The mixture was cooled down to -78°C and lithlumacetylid-ethylendiamin-complex (250 mg, 2.71 mmol) was added. The reaction mixture was stirred for 5h (-78 \rightarrow 20°C).Water (4 mL) was added. The mixture was extracted with dichloromethane (20 mL). The organic phase was dried (Na₂SO₄) and concentrated. Com-

(CDCi3): 5 7.4 (5H, 5 x HC=), 5.0 (2H, CH₂), 2.6 (1H, CH), 1.0 (14H, 2 x CH, 2 x pound 14a (100 mg, 41%) was obtained after flash chromatography. 1H-NMR

Example 57: 5-{[Diisopropyl-(2-methylene-pent-3-enyloxy)-silanyl}-ethynyl}-1-(4-hydroxy-5-(O-triphosphatehydroxymethyl)-tetrahydrofuran-2-yl)-1Hpyrimidine-2,4-dione (compound XIV) ಬ

5-lodo-dUTP (200 mg, 0.56 mmol), diisopropylethylamine (0.1 mL) and 14a (100 lution for 5 min. Tetrakispalladium (57 mg, 0.49 mmol) and Cul (19 mg, 0.1 mmol) were added and the mixture was stirred at 50°C for 5h. Solvent was evaporated off mg, 0.41 mmol) were dissolved in DMF (2 mL). Argon was bubbled through the soand 14b was purified by flash-chromatography. A NMR spectrum revealed the syrup

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consisted of 66%. The syrup was (40 mg) was converted to the corresponding compound VII. TLC indicated full conversion of 14b. Selected NMR data for LH8061a: 14-NMR (MeOD-D₃): 5: 8.3 (1H, HC=), 7.3 (5h, HC=), 6.2 (1H, CH), 5.0 (2H, CH₂), 4.3 (1H, CH), 3.8-3.2 (3H, CH₂, CH), 2.3 (1H, CH₂), 2.2 (1H, CH₂), 1.0 triphosphate (Compound XIV) using the procedure described for the synthesis of

Examples 58 to 63: Preparation of the mononucleotide building block (XV)

(14H, 2 x CH, 4 x CH₃).

Building block XV may be prepared according to the general scheme shown below: 9

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COMPOUND 15a

To a solution of 3-amino-butyric acid (2.06 g, 20 mmol) in NaHCO₃ (50% sat. aq, 25 mL) were added di-tert-butyl dicarbonate (4,36 g, 20 mmol) and acetonitrile (30 mL).

10 The reaction mixture was stirred at room temperature for 18 h. Di-tert-butyl dicarbonate (4,36 g, 20 mmol) was added and the reaction mixture was stirred at room temperature for 18 h.

EtOAc (100 mL) was added and pH was adjusted to 4-5 by addition of NaH₂PO₄. The product was extracted into EtOAc (3 x 100 mL), dried (Na₂SO₄), and evaporated to dryness under vacuum to afford crude product 4.6 g (113%).

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Example 59: Preparation of compound 15b

COMPOUND 15b

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Compound 28 (1,023 g, 5.0 mmol), 3-Eithynyl-phenole (Lancaster, 0.675 g, 12 mmol) and 4-dimethylamino-pyridin (DMAP, 300 mg, 2.5 mmol) were dissolved in EIOAc (10 mL). Dicyclohexyl-carbodiimide (DCC, 2.06 g, 10 mmol) was added to the solution and after 16 h of stirring at room temperature, the reaction mixture was filtered and evaporated to dryness under vacuum. The crude product was purified by silica column chromatography eluting with EtOAc:Heptane gradient (1:3)- (1:2)(v/v). Product yield 720 mg, 73%.

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30 'H NMR (CDCI₃) δ 7.36-7.09 (4H, m, Ph), 4.89 (1H, bs, NH), 4.22 (1H, bm,CH), 3.10 (1H, s), 2.77 (2H, d), 1.40 (3H, 1), 1.32 (3H, d).

Example 60: Preparation of compound 15c

COMPOUND 15c

A solution of 5-lodo-2'-deoxyuridine 3',5'-Di-tert-bulyldimethylsilyl ether (730 mg,

mmol) in anhydrous DMF (3 mL) was stirred at room temperature. N2 was passed 1.25 mmol), triethylamine (250 mg, 2.5 mmol) and compound(15b) (456 mg, 1.5 through the solution for 20 min. 9

Tetrakis(triphenylphosphine)palladium(0) (109 mg, 0.094 mmol) and copper(I) iodide (36 mg, 0.188 mmol) were added and the reaction mixture was stirred at room temperature for 3 d.

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dd, 1'-H), 4.9 (1H, bs), 4.45 (1H, dl), 4,80 (2H, s, CH₂), 4,2 (1H, m), 4.02 (1H, m, 4'-The reaction mixture was evaporated and purified by silica column chromatography 14 NMR (CDCl₃) 8 8.38 (1H, s), 8.08 (1H, s, 6-H), 7.39-7.1 (4H, m, Ph), 6.33 (1H, H), 3.95 (1H, dd, 5'-H), 3.79 (1H, dd, 5"-H), 2,78 (2H, d), 2.36 (1H, m, 2'-H), 2.07 eluting with EtOAc:Heptane gradient (1:3)-(1:2)(v/v). Product yield 807 mg, 85%. (1H, m, 2"-H), 1.46 (9H, s, Bu), 0.93 (9H, s, Bu), 0.91 (9H, s, Bu), 0.15 (3H, s, CH₃), 0.13 (3H, s, CH₃), 0.11 (3H, s, CH₃), 0.09 (3H, s, CH₃).

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Example 61: Preparation of compound 15d

COMPOUND 15d 2

mmol) and tetrabutylammonium fluoride trihydrate (TBAF) (2.36 g, 7.5 mmol) in 20 A solution of compound (15c) (807 mg, 1.06 mmol), glacial acetic acid (1.0 g, 16 mL dry THF was stirred at room temperature for 3 d. The reaction mixture was evaporated and purified by silica column chromatography sluting with (DCM):(MeOH) (9:1) (v/v). Product yield 408 mg, 72%. 2

m, Ph), 6.75 (1H, bd), 6.27 (1H, dd, 1'-H), 4.44 (1H, dt, 4'-H), 3.96 (1H, t, 3'-H), 3.86 H NMR (CD₃OD) § 8.46 (1H, s, 6·H), 7.39 (2H, m, Ph), 7.28 (1H, m, Ph), 7.12 (1H,

(1H, dd, 5'-H), 3.77 (1H, dd, 5"-H), 2,72 (2H, d), 2.35-2.27 (2H, m, 2', 2"-H), 1.46 (9H, s, 'Bu), 1.27 (3H, d). 15

Example 62: Preparation of compound 15e

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COMPOUND 15e

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trimethylphosphate was added (400 µL stock solution (120 mg/mL), 310 µmol). The Compound (15d) (138.5 mg, 260 μmol) was dissolved in 500 μL dry trimethylphosphate. After cooling to 0 °C, a solution of phosphorus oxychloride (POCIs) in dry reaction mixture was stirred at 0 °C for 2h.

Subsequently a solution of tributy/ammoniumpyrophosphate (200 mg, 420 µmol in 1.00 mL dry DMF) and tributylamine (123 mg, 670 µmol in 500 µL dry DMF) was added at 0 °C. The reaction was stirred at room temperature for 3 min. and then stopped by addition of 1 mL 1.0 M triethylammoniumhydrogencarbonate.

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Example 63: Preparation of compound XV 9

COMPOUND XV 5 Removal of N-Boc protection group.

Following phosphorylation, 50 µl of the phosphorylation reaction mixture is adjusted to pH = 1 using HCI and incubated at room temperature for 30 minutes. The mixture is adjusted to pH 5.5 using equimolar NaOH and Na-acetate (pH 5.5) before purification on TLC.

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From the crude mixture, 20 samples of 2 μ l were spotted on kieselgel 60 F_{254} TLC Purification of nucleotide derivatives using thin-layer chromatography (TLC)

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was resuspended in 50-100 µl H₂O to a final concentration of 1-3 mM. The concenrom the nucleotides derivatives using 100% methanol as running solution. Subseshadowing. Kiesel containing the nucleotide-derivative was isolated and extracted quently, the TLC plate is air-dried and the nucleotide-derivative identified by UVcentrifugation and the supernatant was dried in vacuo. The nucleotide derivative wice using 10 mM Na-acetate (pH = 5.5) as solvent. Kieselgel was removed by (Merck). Organic solvents and non-phosphorylated nucleosides were separated

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tration of each nucleotide derivative was evaluated by UV-absorption prior to use in polymerase extension reactions.

Example 64: Polymerase incorporation of different nucleotide derivatives.

- nealed to a template primer using 0.1 and 3 pmol respectively in an extension buffer (20 mM Hepes, 40 mM KCl, 8 mM MgCls, pH 7.4, 10 mM DTT) by heating to 80 °C Different extension primers were 5'-labeled with 32P using T4 polynucleotide kinase for 2 min. and then slowly cooling to about 20 °C. The wild type nucleotide or nuusing standard protocol (Promega, cat# 4103). These extension primers was an-
- samples were mixed with formamide dye and run on a 10% urea polyacrylamide gel film). The incorporation can be identified by the different mobility shift for the nucleotide derivatives compared to the wild type nucleotide. Figure 49 shows incorporation cleotide derivatives was then added (about 100 µM) and incorporated using 5 units of various nucleotide derivates. In lane 1-5 the extension primer 5-GCT ACT GGC ATC GGT-3' was used together with the template primer 5'-GCT GTC TGC AAG AMV Reverse Transcriptase (Promega, part# 9PIM510) at 30 °C for 1 hour. The electrophoresis. The gel was developed using autoradiography (Kodak, BioMax 5 9
- Compound XI; lane 3, Compound IX; lane 4, Compound I; lane 5, Compound II; lane AAG TGA TGA CCG ATG CCA GTA GC-3', and in lane 12-15 the extension primer 5'-GCT ACT GGC ATC GGT-3' was used together with the template primer 5'-GCT GTC TGC AAG TGA CGT AAC CGA TGC CAG TAG C-3'. Lane 1, dATP; lane 2. rGA TAA CCG ATG CCA GTA GC-3', in lane 6-11 extension primer 5'-GCT ACT 6, no nucleotide; lane 7, dCTP; lane 8, Compound VII; lane 9, Compound X; lane GGC ATC GGT-3' was used together with the template primer 5'-GCT GTC TGC 8
- dCTP using different linkers and functional entities. Other polymerases such as Taq, he possibility to incorporate a variety of nucleotide derivatives of dATP, dTTP and ane 14, dTTP and dATP; lane 15, dTTP and Compound X. These results illustrate 10, Compound IV; lane 11, Compound III; lane 12, no nucleotide; lane 13, dTTP; M-MLV and HIV have also been tested with positive results. 22

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Example 65: Polymerase incorporation and hydrolysis of nucleotide derivatives containing cleavable ester linkers.

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ration of various nucleotide derivatives. Lane 1, compound III and Compound II; lane extension primer was annealed with a template primer (5'-TAA GAC CGA TGC CAG one nucleotide derivative can function as the attachment point (non-cleavable linker) and at the same time liberate (cleavable linker) other incorporated nucleotide derivamolar HCI and NaoAc (pH 6.5) and purified by micro-spin gel filtration (BioRad). The film). The incorporation can be identified by the different mobility shift for the nucleoand the incorporated compound III nucleotide derivative with no ester linker is intact. samples were mixed with formamide dye and run on a 10% urea polyacryłamide gel that these nucleotide derivatives can be incorporated by the polymerase in this spe-An extension primer (5'-GCT ACT GGC ATC GGT-3') was 5'-labeled with 32P pound II; lane 4, hydrolysis of compound III and two compound II. The results show derivatives with an ester linker can specifically be hydrolysed on the DNA template using T4 polynucleotide kinase using standard protocol (Promega, cat# 4103) This lives was then added (about 100 µM) and incorporated using 5 units AMV Reverse was treated with 0.1 M NaOH at 50 °C for about 15 min. and then titrated with equitide derivatives compared to the wild type nucleotide. Figure 50 shows the incorpo-IAG C-3') using 0.1 and 3 pmol respectively in an extension buffer (20 mM Hepes, perimental data shows that nucleotide derivatives with linkers containing cleavable 2, compound III and two compound II; lane 3, hydrolysis of compound III and com-Transcriptase (Promega, part# 9PIM510) at 30 °C for 1 hour. Hydrolysed samples cific order. It also shows that one or both the incorporated compound II nucleotide ester can be inserted by the polymerase without reaction with amines in the active 40 mM KCI, 8 mM MgCl₂, pH 7.4, 10 mM DTT) by heating to 80 °C for 2 min. and This illustrates the possibility to incorporate different nucleotide derivatives where then slowly cooling to about 20 °C. The wild type nucleotide or nucleotide derivatives form the DNA template to create a displaying molecule. In addition, this exelectrophoresis. The gel was developed using autoradiography (Kodak, BioMax site of the polymerase or become hydrolysed during the incorporation process.

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Example 66: Polymerase incorporation and cross-linking of nucleotide deriva-

An extension primer (5'-GCT ACT GGC ATC GGT-3') was 5'-labeled with 32P extension primer was annealed with a template primer (5'-TAG ACC GAT GCC AGT slowly cooled to about 20 °C. The nucleotide derivatives was then added (about 100 using T4 polynucleotide kinase using standard protocol (Promega, cat# 4103). This mM KCI, 8 mM MgCl₂, pH 7.4, 10 mM DTT) by heating to 80 °C for 2 min. and then AGC) using 0.1 and 3 pmol respectively in the extension buffer (20 mM Hepes, 40 µМ) and incorporated using 5 units AMV Reverse Transcriptase (Promega, part# S

9PIM510) at 30 °C for 1 hour. The oligonucleotides were then purified using micro-[Bis(sulfonylsuccinimide)suberate] (Pierce, cat# 21580) for about 1 hour at 30 °C. The samples were mixed with formamide dye and run on a 10% urea polyacrylamide gel electrophoresis. The gel was developed using autoradiography (Kodak, spin gel filtration (BioRad). Cross-linking was performed using 10 mM BS₃ 9

show that these nucleotide derivatives can be Incorporated by the polymerase in this BioMax film). Figure 51A and 51B shows the incorporation and cross-linking (CL) of ane 2, cross-linked compound III and compound II. Figure 51B: Lane 1, compound various nucleotide derivatives. Figure 51A: Lane 1, compound III and compound II; III and compound I; Iane 2, cross-linked compound III and compound I. The results specific order. It also shows that compound III, compound II and compound I is 5

antly, the amide groups of the nucleotide derivatives in the major groove are seleccompound II and compound III-compound I mediated by the DNA template. Impormodified by the cross-linking reagent BS₃ (mobility shift) and thereby permit crossinking (CL) between reactive groups on the nucleotide derivatives compound IIIively accessible for modifications which promote cross-linking between different

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Example 67: Polymerase incorporation of various nucleotide derivatives.

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incorporated nucleotide derivatives on the DNA template.

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(20 mM Hepes, 40 mM KCI, 8 mM MgCl₂, pH 7.4, 10 mM DTT) by heating to 80 °C GAT GCC AGT AGC-5') using 0.1 and 3 pmol respectively in the extension buffer An extension primer (5'-TCC GCT ACT GGC ATC GGT-3') was 5'-labeled 4103). This extension primer was annealed with a template primer (5'-TGA ACC with 32P using T4 polynucleotide kinase using standard protocol (Promega, cat#

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Example 68: Polymerase incorporation of various nucleotide derivatives.

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dCTP, dTTP and dATP; lane 2, compound II, compound III and compound XIII. The rivatives after each other using a polymerase. Consequently, the polymerase allows for 2 min. and then slowly cooled to about 20 °C. This template primer was 3'Biotin-(20 mM Hepes, 40 mM KCI, 8 mM MgCl₂, pH 7.4, 10 mM DTT) by heating to 80 °C C6-labeled to prevent extension. The nucleotide derivatives was then added (about and run on a 10% urea polyacrylamide gel electrophoresis. The gel was developed various nucleotide derivatives compared to wild type nucleotides. Lane 1, wild-type various nucleotide derivatives simultaneously in the active site without a significant results show that it is possible to incorporate at least three different nucleotide depart# 9PIM510) at 30 °C for 1 hour. The samples were mixed with formamide dye GAT GCC AGT AGC-3') using 0.1 and 3 pmol respectively in the extension buffer using autoradiography (Kodak, BioMax film). Figure 53 show the incorporation of An extension primer (5'-TCC GCT ACT GGC ATC GGT-3') was 5'-labeled 4103). This extension primer was annealed with a template primer (5'-TGA ACC with 32P using T4 polynucleotide kinase using standard protocol (Promega, cat# 100 µM) and incorporated using 5 units AMV Reverse Transcriptase (Promega, reduction of the catalytic activity.

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Examples 69 to 74; Preparation of polymerase mediated templated molecules

Example 69 Crosslinking of encoded amino groups by urea-bond formation.

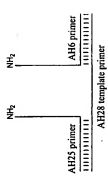
- cleotide kinase using standard protocol (Promega, cat# 4103) and purified by micro-C6-NH2, (Glen research, cat #10-1039-90) was 5'4abeted with 3P using T4 polynuspin gelfiltration. This primer (0.1 pmol) and 2 pmol of a second primer (5'XCA CTT A primer (5'-TCC GCT ACT GGT ATC GGX-3') where X denotes deoxy-thymidine-GCA GAC AGC- 3') were co-annealed with 1 pmol template primer (5'-GCT GTC rGC AAG TGA CCG ATG CCA GTA GC-3") in a hybridisation-buffer (20 mM ഹ
- Hepes, 200 mM NaCl, pH 7.5) by heating to 80 °C for 2 min. and then slowly cooled were mixed with formamide dye and run on a 10% urea polyacrylamide gel electro-Aldrich) was added and the samples incubated at 30 °C for 2 hours. The samples phoresis. The gel was developed using autoradiography (Kodak, BioMax film). A to about 20 °C. Subsequently, 10 mM of N', N'- CarbonylDiimidazole (Sigmaschematic description of this experiment is shown below: . 2 5

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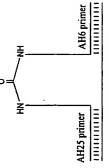
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Cross-linking by urea-bond formation







AH28 template primer

The results shows that adjacent NH2-groups can form a covalent urea-bond by the reaction with CDI. No reaction is observed in the absence of a template sequence which shows that the reaction is dependent on the close proximity of NH2-groups guided by the template sequence. Urea bond formation was also observed when 0.5% formaldehyde was used as cross-linking reagent (data not shown).

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Example 70: Formation of amide bonds by a "fill-in" reaction using a di-amino

nobutane. A primer (5'-TCC GCT ACT GGT ATC GGY-3') where Y denotes deoxy-In this experiment DNA-encoded Carboxylic acids are cross-linked by a 1,4 diami-

- primer (5'-GCT GTC TGC AAG TGA CCG ATG CCA GTA GC-3') in a hybridisationthymidine-C2-COOH (Glen research, cat #10-1035-90), was 5'-labeled with 32P usprimer (5'YCA CTT GCA GAC AGC-3') were co-annealed with 1 pmol template ing T4 polynucleotide kinase using standard protocol (Promega, cat# 4103) and purified by microspin gelfiltration. This primer (0.1 pmol) and 2 pmol of a second
- buffer (20 mM Hepes, 200 mM NaCl, pH 7.5) by heating to 80 °C for 2 min. and then slowly cooled to about 20 °C. Subsequently, 100 mM EDC (Sigma-Aldrich), 10 mM N-hydroxysuccinimide (NHS, Sigma-Aldrich) and 10 mM 1,4 diaminobutane (Merck) mixed with formamide dye and run on a 10% urea polyacrylamide gel electrophoresis. The gel was developed using autoradiography (Kodak, BioMax film). A schewas added and the samples incubated at 30 °C for 2 hours. The samples were 2 13

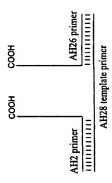
matic description of this experiment is below:

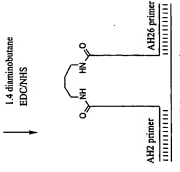
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Cross-linking by "fill-in" reaction





AH28 template primer

proximity of COOH-groups provided by the template sequence. Similar results were functional linker upon formation of amide bonds. No reaction is observed in the ab-The results show that encoded COOH-groups can be covalently coupled by a bisence of a template sequence which shows that the reaction is governed by the obtained using other diamino-linkers such as 1.6 diaminohexane, spermine and spermidine (data not shown).

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Example 71: Polymerase incorporation of nucleotide derivatives and cross-

linking to templated anchor-points.

sion primer was annealed with a template primer (5'-GCT GTC TGC AAG TGA TAA T4 polynucleotide kinase using standard protocol (Promega, cat# 4103). This exten-An extension primer (5'-GCT ACT GGC ATC GGT-3') was 5'-labeled with 32P using buffer (20 mM Hepes, 40 mM KCI, 8 mM MgCl₂, pH 7.4, 10 mM DTT) by heating to was then added (about 100 µM) and incorporated using 5 units AMV Reverse Tran-80 °C for 2 min. and then slowly cooled to about 20 °C. The nucleotide derivatives CCG ATG CCA GTA GC-3') using 0.1 and 3 pmol respectively in the extension Ŋ

scriptase (Promega, part# 9PIM510) at 30 °C for 1 hour. The oligonucleotide complexes were then purified using micro-spin gel filtration (BioRad). A second primer (5-YCA CTT GCA GAC AGC-3') where Y denotes the anchor-point reactive group composition was adjusted to 20 mM HEPES-KOH, 200 mM NaCI, pH=7,5 . Crossdeoxythymidine-C2-COOH, was annealed to the extension complex. The buffer 5

about 2 hours at 30 °C. Relevant samples were subjected to alkaline hydrolysis (0.1 M NaOH, 50 °C for 15 minutes). The samples were mixed with formamide dye and run on a denaturing 10% urea polyacrylamide gel. The gel was developed using autoradiography (Kodak, BioMax film). A schematic outline of this experiment is linking was performed using 100 mM EDC and 10 mM N-hydroxysuccinimid for

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shown below: ಜ

Linking by direct coupling and translocation of a \(\theta\)-Amino acid

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The results show that a reactive group from a nucleotide derivative incorporated by a polymerase can be cross-linked to an anchor point reactive group by a "fill-in" reaction forming amide bonds. Furthermore, the ester linker of the nucleotide derivative is specifically cleaved which allows for the transfer of a templated functional

entity to a templated second entity (anchor point).

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Example 72: Polymerase incorporation of nucleotide derivatives and crosslinking to a templated anchor-point by a "fill-in" reaction.

- An extension primer (5'-GCT ACT GGC ATC GGT-3') was 5'-labeled with ³²P using T4 polynucleotide kinase using standard protocol (Promega, cat# 4103). This extension primer was annealed with a template primer (5'-GCT GTC TGC AAG TGA TAA CCG ATG CCA GTA GC-3') using 0.1 and 3 pmol respectively in the extension buffer (20 mM Hepes, 40 mM KC), 8 mM MgCl₂, pH 7.4, 10 mM DTT) by heating to
- 15 80 °C for 2 min, and then slowly cooled to about 20 °C. The compound II (nucleotide derivative) was then added (about 100 µM) and incorporated using 5 units AMV Reverse Transcriptase (Promega, part# 9PIM510) at 30 °C for 1 hour. The oligonucleotide complexes were then purified using micro-spin gel filtration (BioRad). A second primer (5-XCA CTT GCA GAC AGC-3') where X denotes the anchor-point reactive
- group deoxythymidine-C6-NH₂, was annealed to the extension complex. Cross-linking was performed using 10 mM BS, [Bis(sulfonylsuccinimide)suberate] (Pierce, cat# 21580) for about 2 hours at 30 °C. Relevant samples were subjected to alkaline hydrolysis (0.1 M NaOH, 50 °C for 15 minutes). The samples were mixed with formamide dye and run on a denaturing 10% urea polyacrylamide gel. The gel was
- 25 developed using autoradiography (Kodak, BioMax film). A schematic outline of this experiment is shown below:

Linking by "fill-in" and B-Amino acid translocation

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A copy of the gel is shown in Figure 54. Lane 1: no nucleotides, lane 2: dTTP, lane followed by alkaline hydrolysis, lane 6: dTTP followed by BS3 cross-linking, lane 7: 3: compound I, lane 4: dTTP followed by alkaline hydrolysis, lane 5: compound I

inking and alkaline hydrolysis, and lane 9: compound I followed by BS3 cross-linking active group by a "fill-in" reaction forming amide bonds. Furthermore, the ester linker of the nucleotide derivative is specifically cleaved which allows for the transfer of a derivative incorporated by a polymerase can be cross-linked to an anchor point reand alkaline hydrolysis. The results show that a reactive group from a nucleotide compound I followed by BS₃ cross-linking, lane 8: dTTP followed by BS₃ crossemplated functional entity to a templated second entity (anchor point). ß 9

Example 73: Polymerase incorporation of two nucleotide derivatives and the cross-linking between 3 encoded entities.

and X was then added (about 100 µM) and incorporated using 5 units AMV Reverse sion primer was annealed with a template primer (5'-GCT GTC TGC AAG TGA GTA An extension primer (5'-GCT ACT GGC ATC GGT-3') was 5'-labeled with 32P using Г4 polynucleotide kinase using standard protocol (Promega, cat# 4103). This extenbuffer (20 mM Hepes, 40 mM KCI, 8 mM MgCl₂, pH 7.4, 10 mM DTT) by heating to 30 °C for 2 min, and then slowly cooled to about 20 °C. The nucleotide derivative V primer (5-YCA CTT GCA GAC AGC-3') where Y denotes the anchor-point reactive franscriptase (Promega, part# 9PIM510) at 30 °C for 1 hour. The oligonucleotide group deoxythymidine-C2-COOH, was annealed to the extension complex. The CCG ATG CCA GTA GC-3') using 0.1 and 3 pmol respectively in the extension complexes were then purified using micro-spln gel filtration (BioRad). A second 15 ೪ 32

NaOH 50 °C, 15 minutes). Formamide dye was added to the samples before loading before addition of 100 mM EDC and 10 mM N-hydroxysuccinimid. This results in the on a 10 % Urea polyacrylamide gel. The gel was developed using autoradiography cross-linking of NH2-groups of MG91 and the COOH group of V and the COOH of buffer composition was adjusted to 20 mM HEPES-KOH, 200 mM NaCl, pH = 7.5 the second primer. Suitable samples were subjected to alkaline hydrolysis (0.1 M (Kodak, BioMax film). A schematic representation of this experiment is shown be-ဓ္က

Linking of 3 encoded functional entities

Template

Template

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This result shows that three encoded functional entities can be cross-linked. Furthermore, a specific linker can be selectively cleaved.

Example 74: Polymerase incorporation and 0-amino acid translocation ("Zip-

plng").

An extension primer (5'-GCT ACT GGC ATC GGT-3') was 5'-labeled with ³²P using T4 polynucleotide kinase using standard protocol (Promega, cat# 4103). This extension primer was annealed with a template primer (5'-TAG ACC GAT GCC AGT AGC) using 0.1 and 3 pmol respectively in the extension buffer (20 mM Hepes, 40

- slowly cooled to about 20 °C. The nucleotide derivatings to 80 °C for 2 min. and then slowly cooled to about 20 °C. The nucleotide derivatives II and III was then added (about 100 µM) and incorporated using 5 units AMV Reverse Transcriptase (Promega, part# 9PIM510) at 30 °C for 1 hour. The oligonucleotides were then punified using micro-spin gel filtration (BioRad) followed by lyophilization. The oligonucleotides
- cleotide complex was dissolved in pyridine and Scandiumtriflourmethanesulphonate (catalyst) in pyridine was added to a final concentration of 10 mM and the reaction mixture incubated at 50 °C for 1 hour. Relevant samples were subjected to alkaline hydrolysis using 0.1 M NaOH at 50 °C for 15 min. Formamide dye was added to the samples before loading on a 10 % Urea polyacrylamide gel. The gel was developed using autoradiography (Kodak, BioMax film). A schematic representation is shown

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Zipping and translocation of a β–Amino acld

The results show that a reactive group of a functional entity can react with a reactive group of an other functional entity forming an amide bond. The reaction results in a translocation of a functional entity onto a second functional entity with simultaneous cleavage of the linker connecting the first functional entity and the nucleotide derivative that encode said functional entity. In this experimental set-up a di-peptide comprising two β-amino acids is produced. Thus, incorporation on a DNA template of several (3 or more) nucleotide derivatives comprising β-amino-acids as functional entities would allow multiple translocation events producing β-peptides acids comprising 3 or more β-amino acids. In this example the reaction between functional entity reactive groups occurs in non-aqueous environment. In a preferred aspect the

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reaction between functional entity reactive groups occurs directly upon incorporation of a nucleotide derivative comprising said function entity by a "zipping" reaction. This can be accomplished by increasing the reactivity of the ester linkage by introducing various chemical entities such as thioesters, phenolic esters, thiophenolic esters, ditri- or tetra-fluoro-activated phenolic- or thiophenolic esters or N-hydroxysuccimide

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Example 75; in silico experiment A structural description of a template-

displayed molecule created using polymerase incorporation of nucleotide

derivatives.

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One aspect of the present invention utilizes a suitable polymerase for specific incorporation of nucleotide derivatives on a DNA template. This incorporation is accomplished using a template containing coding elements. The template is utilized by the polymerase to incorporate the nucleotide derivatives in a specific order based on these coding elements (Figure 55). This process is specific due to the recognition groups in the nucleotide derivatives.

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shown in Figure. 55A. The consecutive incorporation of the nucleotide derivatives by entities in or outside the major groove of the DNA strand exposed to the solvent as the nucleotide derivatives. The linker design will differ dependent on which type and each functional entity (examples shown in Figure 11-21). In addition, the DNA tem-The different nucleotides are modified at specific positions (e.g. Figure 9) to permit incorporation by the polymerase and at the same time expose the linked functional olate will arrange the functional entities in specific geometry dependent on the helical structure of the DNA template. This geometry can for example be controlled by how the reactive groups are arranged in the functional entities. The linker can also different types of linkers that join the functional entity and complementing element. Thus, the linker is designed to favour the reaction between the reactive groups on the polymerase will allow various reactions to occur between the linked functional antities. The reactions are determined by the type of reactive groups integrated in be designed to guide the reaction between the reactive groups in a specific direcion. Various reactive groups can also be used to direct the reaction between the ಜ 22 ജ

reactive groups. The close proximity and the optimized geometry of the nucleotide

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derivatives will drastically enhance the reaction rate between the reactive groups in the different functional entities. The reaction rate between the reactive groups is fast due the high local concentration of the incorporated nucleotide derivatives on the DNA template molecule compared to if they were allowed to diffuse freely in solu-

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Figure 55A shows one example where nucleotide derivatives Compound II, compound X and compound V are incorporated by a polymerase after each other on the same DNA template. The synthesis of these nucleotide derivatives are described in detail shows. The experimental data showing AMV Deuces transcriptuse incorpora-

detail above. The experimental data showing AMV Reverse transcriptase incorporation of these nucleotide derivatives can be seen in example 64. These incorporated nucleotide derivatives are structurally arranged, by the linker connecting the complementing element and the functional entity, to promote reaction between the reactive groups on each nucleotide derivatives. The distance between the amine in

compound II and the COMPOUND X amine in the long side chain is calculated to be between 3.1 A and 17.5 A and the distance between the amine in compound II and the COMPOUND X amine in the short side chain to be between 3.0 A and 14.6 A dependent on the precise orientation of the linker and the functional entity on the DNA template. The distance between the carbonyl carbon in nucleotide derivative compound V and the long side chain amine in nucleotide derivative COMPOUND X is between 4.2 A and 19.8 A and the distance to the short side chain COMPOUND X amine is calculated to be between 3.7 A and 16.5 A also dependent of the precise orientation. The close proximity of the nucleotide derivatives compound II, COMPOUND X and 1973 on the DNA template will promote a chemical linkage of the

25 reactive groups in these nucleotide derivatives.

These three nucleotide derivatives can be linked together through their reactive groups using various chemical reagents. One possible reagent to use is BS₃ [Bis(sulfony)succinimide)suberate] (Pierce, cat# 21580). Typically a concentration of about 0.25 – 10 mM is used of this analog. This reagent will cross-link two amines between nucleotide derivatives compound II and COMPOUND X. This particular reagent will insert a spacer of eight carbons between the reactive groups and is capable of bridging a distance of 11.2 A in the extended conformation. Thus, the BS₃ linker is capable of linking the amines of compound II and either of the amines of compound X. There are other reagents that could be used (longer or shorter) to

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obtain almost any type of spacers between the reactive amine groups. The carbox-ylic acid on nucleotide derivative compound V and one of the amines on nucleotide derivative COMPOUND X can be linked together using for example 1-Ethyl-3-(3-dimethylaminopropyl) carbodilmide (EDC) and N-Hydroxysuccinimide (NHS). This

- foreaction will make a direct connection between the reactive groups on nucleotide derivatives COMPOUND X and compound V. These two reactions result in a new molecule composed of these nucleotide derivatives covalently attached to each other through the coupling reagents (Fig. 55B). This particular DNA templatemediated molecule is produced using both fill-in (BS₃) and direct coupling
- (EDC/NHS) chemistry. Examples of cross-linking between incorporated nucleotide derivatives are shown above. Other types of coupling approaches that could be used are zipping by translocation or ring opening. These coupling strategies need other types of linker design as described in this invention.
- At this stage, all the functional entities are still attached to the DNA template through the linker joining the functional entity and the complementing element. The ester element integrated in the linker of nucleotide derivatives compound II and COM-POUND X can specifically be hydrolysed (see example 65 for experimental details) to liberate the functional entities of these two nucleotide derivatives from the DNA
- template. This hydrolysis reaction results in a new molecule that is only attached to the DNA template through the linker in the compound V nucleotide derivative (fig. 55C). This molecule can then extend out from the DNA template into the solution and become accessible (displayed) for interaction with other molecules in the sotution.

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This templated molecule, as part of a library of many different templated molecules, can finally be used in a selection procedure to identify molecules that bind to various targets. A detailed description of the selection procedure can be found elsewhere herein.

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Example (Model) 76; PNA synthesis - base linked

PNA monomers are linked to complementing elements via cleavable benzyl- or benzyloxycarbonyl moieties bound to the base part of each PNA monomer. A carboxylic

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acid is used as anchor point to the oligonucleotide complex. Each building block is annealed to a oligonucleotide template (not shown).

Step A: Polymerization

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To an aqueous buffered solution (10uL, 1M NaCl, 100-500 mM buffer pH 6-10, preferably 7-9) of oligonucleotide complexes (0.1-100 uM, preferably 0.5-10 uM) is added a peptide coupling reagent (0.1 mM – 100 mM, preferably 1-10 mM) exemplified by but not limited to EDC, DCC, DIC, HATU, HBTU, PyBoP, PyBoP or N-methyl-2-chloropyridinium tetrafluoroborate and a peptide coupling modifier (0.1 mM-1 uM, preferably 1-10 mM) exemplified by but not limited to NHS, sulpho-NHS, HOBt, HOAt, or DhbtOH in a suitable solvent (1uL) e.g. water, methanol, ethanol, dimethylformamide, dimethylsulfoxide, ethylene glycol, acetonitrile or a mixture of these. Reactions run at temperatures between -20 °C and 60 °C. Reaction times are between 1h and 1 week, preferably 1h-24h.

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The above procedure exemplifies the polymerisation on an 11 uL scale, but any 15 other reaction volume between 1.1 uL and 1.1L may be employed.

Step B: Linker cleavage and deprotection

Cbz- and Benzyl protective groups may be removed by a variety of methods, [Greene;1999;] Due to its mildness, catalytic reduction is often the method of choice. Combining an insoluble hydrogenation catalyst e.g. Pd/A₂O₃. Pd/CaCO₃, Pd/C, PtO₂, or a soluble one e.g. Wilkinsons catalyst and a hydrogen source exemplified but not limited to H₂, ammonium formiate, formic acid, 1,4-cyclohexadien, and cyclohexene in a suitable solvent like water, methanol, ethanol, dimethylformamide, dimethylsulfoxide, ethylene glycol, acetonitrile, acetic acid or a mixture of these with the oligo nucleotide complexes removes the Cbz- and benzyl protective groups.

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Scheme for building block synthesis:

Step A, B:

Upon completion, (6-Aminopurin-9-yl)-acetic acid ethyl ester (5mmol) in pyridine (20 mL) is added and left to react 16h at rt. Volatiles are removed in vacuo and the resiice/water bath is added (4-Ethynylphenyl)methanol (5 mmol) dissolved in DCM (20 mL) dropwise. After 1h the ice bath is removed. The reaction is monitored by TLC. To a DCM solution (20 mL) of 4-nitrophenolchloroformiate (5 mmol) cooled on an due purified by chromatography.

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Step C, D:

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Steps C[Hyrup;1996; Bioorganic & medicinal chemistry, 5-23] and D[Schmidt;1997; Nucleic Acids Research; 4792-4796, Böhler; 1995; Nature;] are known from the literature.

Step E

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volatiles followed by chromatography affords the desired modified nucleoside that is the alkyne (0.69 mmol, 2 eq), DIEA (0.25 mL) is purged with Ar for 5 min. Tetrakis triphenylphosphine palladium (0.03 mmol, 0.1 eq) and Cul (0.07 mmol, 0.2 eq) is added and the mixture is heated to 50 °C and kept there for 20 h. Evaporation of A DMF solution (2 mL) of the protected iodo substituted nucleoside (0.34 mmol),

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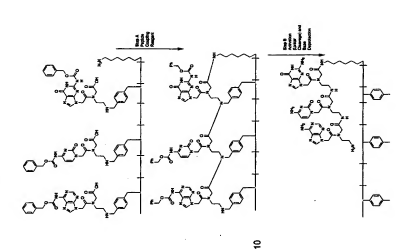
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converted into its corresponding phosphor amidite and incorporated into an oligonucleotide.

Example (model) 77: PNA Synthesis - Nitrogen linked

point to the oligonucleotide complex. Each building block is annealed to a oligonu-PNA monomers are linked to complementing elements via cleavable benzyl moieties bound to the base part of each PNA monomer. An amine is used as anchor cleotide template (not shown). ໝ



Designates a valence bond between modified nucleotides.

Step A: Polymerization

these. Reactions run at temperatures between -20 °C and 60 °C. Reaction times are To an aqueous buffered solution (10uL, 1M NaCI, 100-500 mM buffer pH 6-10, prefadded a peptide coupling reagent (0.1 mM – 100 mM, preferably 1-10 mM) exemplimM-1 uM, preferably 1-10 mM) exemplified by but not limited to NHS, sulpho-NHS, dimethylformamide, dimethylsulfoxide, ethylene glycol, acetonitrile or a mixture of HOBt, HOAt, or DhbtOH in a suitable solvent (1uL) e.g. water, methanol, ethanol, erably 7-9) of oligonuclectide complexes (0.1-100 uM, preferably 0.5-10 uM) is methyl-2-chloropyridinium tetrafluoroborate and a peptide coupling modifier (0.1 fled by but not limited to EDC, DCC, DIC, HATU, HBTU, PyBoP, PyBroP or Nbetween 1h and 1 week, preferably 1h-24h.

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The above procedure exemplifies the polymerisation on a 11 uL scale, but any other reaction volume between 1.1 uL and 1.1L may be employed.

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Step B:

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cyclohexadien, and cyclohexene in a suitable solvent like water, methanol, ethanol, mixture of these with the oligo nucleotide complexes removes the Cbz- and benzyl Pd/CaCO₃, Pd/C, PtO₂, or a soluble one e.g. Wilkinsons catalyst and a hydrogen dimethylformamide, dimethylsulfoxide, ethylen glycol, acetonitril, acetic acid or a method of choice. Combining an insoluble hydrogenation catalyst e.g. Pd/Al₂O₃, source exemplified but not limited to H2, ammonium formiate, formic acid, 1,4-Cbz- and Benzyl protective groups may be removed by a variety of methods, [Greene and Wuts;1999;] Due to its mildness, catalytic reduction is often the protective groups.

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Example 78 (model): Polysaccharides

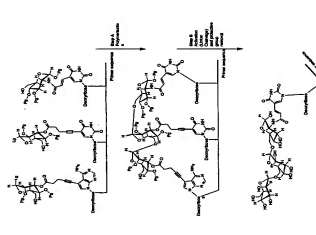
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General scheme for polysaccharide synthesis

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Step A

plate (not shown) and extended with modified nucleotides carrying hexose units. Pg A primer sequence modified with a carboxylic acid (e.g. Glen Research Carboxy-dT cat. No. 10-1035-) that has been attached to a 2-amino-sugar is annealed to a temis a protection group[Seeberger;2000; Chem. Rev.; 4349-

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thenot, phosphate esters and sugar nucleoside phosphates or sugar phosphates for (SiR₃ wherein R is tower alkyl), Lg is a leaving group typical for carbohydrate chemistry exemplified by but not limited to halogen, trichloroacetamidato, mercaptan,

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4393, Seeberger, 2001;] exemplified by but not limited to Ac, Bz, Lev, Piv, Silanes

enzymatic[Wong;1994; Tetrahedron Organic Chemistry Series;] carbohydrate synthesis. Polysaccharides may also be synthesised using glycals.

Step B: Linker Activation

The ester linkages are cleaved with aqueous hydroxide at pH 9-12 at room temperadimethylsulfoxide, ethylene glycol, acetonitrile or a mixture of these. If Pg is Ac or ure, 16 h in a suitable solvent like water, methanol, ethanol, dimethylformamide, other base labile protective group, these are removed as well.

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menting element. This example shows a 1-6 coupled trimer but any combination of Carbohydrates have several OH-functionalities allowing attachment to the complebuilding blocks may be used.

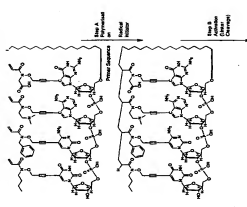
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to fold into secondary structures hence facilitating the synthesis of polysaccharides Attaching carbohydrate units to a template may lessen the tendency of these units

Example (Model) 79: Acrylamide

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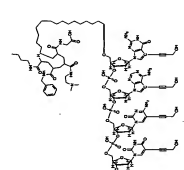
General scheme for a polyacrylamid synthesis:



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oligo nucleotide template (not shown) and extended with modified nucleotides carry-A terminally modified primer sequence carrying an lodine atom is annealed to an ing M-substituted acrylamide units.

Step A: Polymerisation

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Acrylamides are polymerized in a cascade radical reaction starting by abstraction of the iodine atom by a radical initiator forming a carbon atom based radical. To an aqueous buffered solution (10uL, 1M NaCl, 100-500 mM buffer pH 6-10, preferably 7-9) of oligonucleotide complexes (0.1-100 uM, preferably 0.5-10 uM) carryarably 1-10 mM) exemplified by but not limited to peroxymonosulfate, AIBN, di-tert solvent (1uL) e.g. water, methanol, ethanol, dimethylformamide, dimethylsulfoxide, ing M-substituted acrylamld units is added a radical initiator (0.1mM-100mM, prefbutylperoxide, tert butylperoxide, hydrogen peroxide or lead acetate in a suitable 9

ethylene glycol, acetonitrile or a mixture of these, optionally applying UV-light, ultrasound or microwaves. Reactions run at temperatures between -20 °C and 100 °C, preferably between 0 °C and 60 °C. Reaction times are between 1h and 1 week, preferably 1h-24h. ŧ

The above procedure exemplifies the polymerisation on a 11 uL scale, but any other reaction volume between 1.1 uL and 1.1L may be employed

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Step B: Activation

The N-O bond is susceptible to cleavage by reduction using hydrogenation catalysts and a suitable hydrogen source or in the presence of certain metal salts.

between -20 °C and 100 °C, preferably between 0 °C and 60 °C. Reaction times are To an aqueous buffered solution (10uL, 1M NaCl, 100-500 mM buffer pH 4-10, prefadded reductants (0.1mM-100mM, preferably 1-10 mM) exemplified by but not limsolvent (1uL) e.g. water, methanol, ethanol, dimethylformamide, dimethylsulfoxide, ited to samanum(II) iodide, tin(II) chloride or manganese(III) chloride in a suitable ethylene glycol, acetonitrile or a mixture of these. Reactions run at temperatures erably 4-7) of oligonucleotide complexes (0.1-100 uM, preferably 0.5-10 uM) is between 1h and 1 week, preferably 1h-24h.

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The above procedure exemplifies the polymerisation on a 11 uL scale, but any other reaction volume between 1.1 uL and 1.1L may be employed.

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Building block synthesis:

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Tetrakis triphenylphosphine palladium (0.3 mmol, 0.1 eq) and Cul (0.7 mmol, 0.2 eq) A DMF solution (20 mL) of the protected iodo substituted nucleoside (3.4 mmol), the alkyne (6.9 mmol, 2 eq, Aldrich P51338), DIEA (2.5 mL) is purged with Ar for 5 min. is added and the mixture is heated to 50 °C and kept there for 20 h. Upon cooling,

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the mixture is added 700 mL diethylether. The organic phase is washed with ammothat is purified by column chromatography (silica gel, Heptane/Ethyl acetate eluent). lowed by stripping with toluene (400 mL) affords the desired modified nucleoside nium chloride (sat, aq, 250 mL) and water (250 mL). Evaporation of volatiles fol-

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Step B:

To the modified nucleoside obtained in Step A (2 mmol) in ethanol (30 mL) is added reaction is monitored by TLC. Upon completion volatiles are removed in vacuo and hydrazine hydrate (400 mg, 8 mmol, 4 eq.) and the mixture is stirred at 20 °C. The the residue purified by chromatography.

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The amine obtained in Step B (0.5 mmol) is added DMF (10 mL), benzaldehyde (0.6 Reacts at 20 °C, 16 h and is quenched with NaHCO₃ (aq, 10 mL, 5%) and extracted with ethyl acetate (3x100 mL). The combined organic phase is washed with NH4Cl mmol, 1.2 eq), acetic acid (100 uL, 1%) and sodium cyanoborohydride (0.6 mmol). sat, aq, 50 mL) and water (50) mL and dried over Na₂SO₄. Upon evaporation of ethyl acetate, the residue may be purified by chromatography.

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Step D:

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HCI(aq) (0.1 M, 3 mL), NaHCO3 (aq, 3 mL, 5%) and water (3 mL). Upon evaporation educed under vacuum. The residue is taken up in ethyl acetate and is washed with ride (0.15 mmol) in dichloromethane (2 mL) is added dropwise. Upon 1h reaction at 0 °C the temperature is allowed to raise to 20 °C and the reaction is quenched after 1h with NaHCO3 (aq, 3 mL, 5%). The phases are separated and the organic phase of ethyl acetate, the product is stripped with toluene (2x20 mL), purified by chromapresence of 2,6-di tertbutylpyridine (0.4 mmol) and cooled to 0 °C where acrylchlo-The product obtained in Step C (0.1 mmoL) is dissolved in dichloromethane in the ography and converted into the desired building block type, e.g. a 5'-triphosphate.

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Example (Model) 80; Synthesis of B-peptides

General scheme for β-peptide synthesis:

Step A

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The amine precursor may be an amine carrying a protective group[Greene and Wuts;1899;] exemplified by but not limited to benzyl carbamate, paramethoxybenzyl carbamate, 2-Trimethylsilylethyl carbamate, 2,2,2-Trichloroethyl Carbamate. These protective groups are removed by hydrogenolysis, mild acid treatment, fluoride treatment and treatment with Zn dust respectively. Alternatively, the amine precursor may be a nitro group or an azide. Both are converted into amines by reduction. The latter is also reduced under mild conditions using phosphines.

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Step B

The free amine generated in step A attacks the neighbouring NTA unit to start the cascade.

5 Step C

Linker cleavage is carried out using UV radiation (250-500 nm) on a buffered solution of oligonucleotide complexes (pH 5-10) to partially release a beta peptide.

Example (Model) 81: B-peptoid synthesis

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Step B St

Step A

Wuts;1999;] exemplified by but not limited to benzyl carbamate, paramethoxybenzyl treatment and treatment with Zn dust respectively. Alternatively, the amine precursor carbamate, 2-Trimethy/silylethy/ carbamate, 2,2,2-Trichloroethy/ Carbamate. These may be a nitro group or an azide. Both are converted into amines by reduction. The protective groups are removed by hydrogenolysis, mild acid treatment, fluoride The amine precursor may be an amine carrying a protective group[Greene and latter is also reduced under mild conditions using phosphines.

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Step B

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unit initially forming an unstable aminal due to the ring opening. This collapses to an The free amine generated in step A attacks the neighbouring [1,3]Oxazinan-6-one aldehyde releasing a secondary amine which is now able to continue the cascade resulting in this case in a beta peptoid.

Building block synthesis

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Example (Model) 82: Polyamide synthesis

Alternating monomer building blocks of type X-X and Y-Y are incorporated (principle depicted in Figure 16) followed by a polymerization step resulting in bond formation between X and Y on neighbouring monomers.

Building block synthesis ß

Step A: 2+2 cycloaddition

2-Allyl-matonic acid dimethyl ester (1 mmol) and Chlorosulfonyl isocyanate (1 mmol) are mixed in THF at 20 °C and left to react 7 days. The crude product is used with-

out purification.

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Step B: Di-amine protection

1,3-Diamino-propan-2-ol (1 mmol) and trifluoroacetic anhydride (2 mmol) is mixed in diethylether at 0 °C and left to react at this temperature 4h. The reaction mixture is extracted with 1M HCI, NaHCO3 (aq) and water. The product is obtained by evaporation of the organic phase

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Step C: Carbamate formation

(1.5 mmol) and left to react at 20 °C, 16h. The crude product is used without purifi-2,2,2-Trifluoro-N-[2-hydroxy-3-(2,2,2-trifluoro-acetylamino)-propyl]-acetamide (1.5 mmol) obtained in step B is dissolved in THF along with chlorosulfonyl isocyanate 2

Step D: Reductive amination

mmol), sodium cyanoborohydride (6 mmol) and acetic acid. Upon stirring overnight volatiles are removed and the product is purified by crystallisation or chromatogra-The aldehyde (5 mmol) is dissolved in a minimum MeOH and added an amine (6

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Step E: Sulfonamide formation

step D in a water/THF mixture in the presence of base and left to react at 20 °C, 4h. The crude product from step A or step C in THF is added to the amine obtained in Then the mixture is refluxed over night. Upon cooling, the solvent is removed and the residue purified by chromatography.

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Oligo building block preparation

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The protected diamines and diacids are attached to modified oligonucleotides carrying a primary amino functionality using EDC and NHS in an aqueous buffer (pH 5-8, preferably 6-7). The protective groups (both methyl esters and trifluoro acetamides)

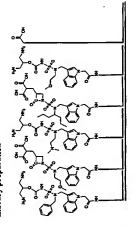
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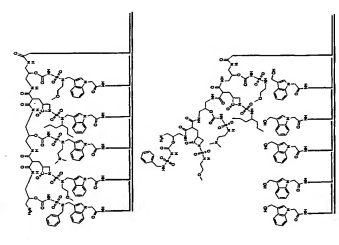
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main on the building blocks and are removed after annealing to the oligonucleotide are removed in aqueous buffer (pH 10-12). Alternatively, the protection groups re-

Library preparation മ





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Polymerisation:

ing di-amines and di-carboxylic acids is added a peptide coupling reagent (0.1 mM -HATU, HBTU, PyBoP, PyBroP or M-methyl-2-chloropyridinium tetrafluoroborate and (1uL) e.g. water, methanol, ethanol, dimethylformamide, dimethylsulfoxide, ethylene glycol, acetonitrile or a mixture of these. Reactions run at temperatures between -20 To an aqueous buffered solution (10uL, 1M NaCI, 100-500 mM buffer pH 6-10, preferably 7-9) of oligonucleotide complexes (0.1-100 uM, preferably 0.5-10 uM) carrya peptide coupling modifier (0.1 mM- 100 mM, preferably 1-10 mM) exemplified by 100 mM, preferably 1-10 mM) exemplified by but not limited to EDC, DCC, DIC, but not limited to NHS, sulpho-NHS, HOBt, HOAt, DhbtOH in a suitable solvent °C and 60 °C. Reaction times are between 1h and 1 week, preferably 1h-24h.

Activation (Linker cleavage):

The above procedure exemplifies the polymerisation on a 11 uL scale, but any other

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reaction volume between 1.1 uL and 1.1L may be employed.

Linkers are cleaved by treatment with acid pH 0-5, at 0-40 °C for 10 min-10 h. 5

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Example (model) 83. Isolation of a-peptide ligand to Glutathione S-transferase (GST) from a library of templated α-peptides.

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A) Nucleotide derivative synthesis

The synthetic strategy for three nucleotide derivatives is shown in the scheme below with a detailed description of the synthesis. Examples of other synthesized α -amino acid nucleotide derivatives can be found in the literature (e.g. Ito et al. (1980) J.

Amer. Chem. Soc. 102: 7535-7541; Norris et al. (1996) J. Amer. Chem. Soc. 118: 5769-5803; Celewicz et al (1998) Pol. J. Chem. 72: 725-734). S

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Synthesis of LH1, LH7: EDC (3.2 mmol) is added to an ice-water cooled solution of either N-{fert-butoxycarbonyl}-lent-butoxy glutamate (3.0 mmol) or N-{fert-butoxycarbonyl}-glycine (3.0 mmol) in dichloromethane (10 mL). A solution of 4-dimethylamino pyridine (0.3 mmol) and 5-hexynol (4.6 mmol) in dichloromethane (1 mL) is added. The reaction mixture is stirred for 1 h. at 0°C, then at room temperature overnight. Solvent was evaporated off and the residue is taken up in diethyl ether. The slurry is washed with HCi (0.1 M, 25 mL), saturated NaHCO₃ (25 mL) and brine (25 mL), then concentrated to oil. The product is purified by flash-chromatography.

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Synthesis of LH4: 6-lodohexyne (6 mmol) and K₂CO₃ (6 mmol) is added to a solution of N-(tert-butoxycarbonyl)-cysteine (3 mmol) in methanol (5 mL) and DMF (5 mL), The reaction mixture is stirred for 1 day at 40°C, then concentrated and worked-up by column chromatography.

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Synthesis of LH2, LH5 and LH8: Tetrakis(triphenylphosphine)palladium (0.6 mmol) and Cul (0.2 mmol) is added to a degassed solution of the lodo nucleoside (1 mmol), the alkyne (2 mmol) and ethyldiisopropyl amine (2 mmol) in DMF or ethanol (4 mL). The reaction mixture is stirred under an atmosphere of argon. The reaction was followed by TLC. The reaction is stirred at 50°C if no reaction occurred at room temperature. The reaction mixture is concentrated to syrup and worked-up by RP-HPLC (eluent: water → methanol). The corresponding (err-butyldimethyl sily) protected iodo nucleoside is used instead of the unprotected nucleiside when the primary hydroxyl group is acylated in the course of the reaction. The silyl ether is cleaved after the Sonogashira coupling by treating the compound with tetrabutyl ammonium fluoride (4 eq.) in a solution of ethanol and acetic acid (8 eq.) for 1 day followed by concentration and work-up by RP-HPLC (eluent: water → methanol).

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Synthesis of LH3, LH6 and LH9: Phosphooxychloride (0.11 mmol) is added to an ice-water cooled solution of the nucleoside (0.1 mmol) in trimethyl phosphate (1 mL). The reaction mixture is stirred under an atmosphere of argon at 0°C for 1h. A solution of bis-n-tributylammonium pyrophosphate (0.2 mmol) in DMF (1 mL) and n-tributylamine (0.3 mmol) is then added. The reaction mixture is stirred for 10 minutes then water (1 mL) was added. The mixture is neutralized with triethylamine and stirred at room temperature for 6 h, then concentrated in vacuo and worked-up by

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ion pair exchange RP HPLC (eluent 100 mM triethylammonium acetate → 100 mM triethylammonium acetate in 80% acetonitrile). Removal of buffer salts from the nucleotide is carried out by adding water (100 µl) to the mixture and then concentrating the slurry at 0.1 mmHg several times finally followed by a gel filtration (eluent: wa-

B) Library design and nucleotide derivative incorporation

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A templated library can be produced by extension of a primer annealed to a template primer. The template primer encodes the library and can be prepared using

- standard procedures, e.g. by organ synthesis with phosphoramidite. To generate various types of oligonucleotide libraries one can for example use redundancies, mixed phosphoramidite or doping in synthesizing the oligonucleotides. These oilgonucleotide libraries can be purchased from a supplier making customer defined oligonucleotides (e.g. DNA Technology A/S, Denmark or TAG Copenhagen A/S,
 - 15 Denmark).

Here, An extension primer (5'-GCT ACT GGC ATC GGT-3') is used together with a template primer (5'-GTA ATT GGA GTG AGC CDD DAC CGA TGC CAG TAG C-3') where D (underlined, using the ambiguity definition from International Union of Biochamiers) is gither A. Got The extension primer is complemented to the construction of the cons

- chemistry) is either A, G or T. The extension primer is complementary to the template primer as shown below. During extension the primer is extended past the DDD-sequence, leading to insertion of T-, C-, or A-nucleotide derivatives at there position, according to the sequence of the individual templates. Upon polymerization of the α-amino and precursors attached to the nucleotides, and cleavage of the
- linker that connect the amino and the nucleotide, a library with a theoretical diversity of at least $3^3=27$ different peptides is created.

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Library design

extension of

B = 1, C or A wild type nucleoides

GCT ACT GGC ATC GGT HHH GGC TCA CAC CAA TTA

GCT ACT GGC ATC GGT HHH GGC TCA CAC CAA TTA CGA TGA CCG TAG CCA DDD CCG AGT GAG GTT The extension primer is annealed with the template primer, using about 3 pmol of each primer in an extension buffer (20 mM Hepes, 40 mM KCI, 8 mM MgCl₂, pH 7.4, 10 mM DTT), by heating to 80 °C for 2 min and then slowly cooling to about 20 °C. The nucleotide derivatives are then added to a concentration of about 200 µM each, and incorporated using 5 units AMV Reverse Transcriptase (Promega, part# 9PIM510) at 30 °C for 1 hour. Unincorporated nucleotide derivatives are removed using a spin-column (BioRad). Further extension may be performed by adding wild type dNTP using the same conditions described for the nucleotide derivatives. Alternatively, an oligonucleotide that anneal to the sequence downstream of the DDD sequence is added prior to the extension. The double stranded product is purified and transferred to another buffer (100 mM Na-phosphate buffer, pH 8.0) using a spin-column (BioRad).

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C) Polymerization and linker cleavage

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The reactive groups of the incorporated nucleotide derivatives are linked together using 1-Ethyl-3-(3-dimethylaminopropyl) carbodilmide (EDC) and N-Hydroxysuccinimide (NHS). This is a routine procedure for covalent coupling amines

and carboxyl groups. Examples of coupling conditions are described in the literature

(e.g. NHS coupling kit, IAsys, code # NHS-2005).

EDC and NHS are added to the purified double stranded extension product at appropriate final concentrations of about 100 mM and 10 mM, respectively. This reac-

tion is incubated at 30 °C for 2-16 hours. Excess linking reagents is removed using a

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spin column. Hydrolysis of hydrolysable linkers is achieved by incubating the sample at pH 11 (e.g. 0.2 M NaOH) for 15 min at 50 °C.

D) Selection

One of the possible templated molecules in this particular fibrary, when using nucleotide derivatives LH3, LH6 and LH9, is glutathione (Glu-Cys-Gly). The incorporation, reaction between the reactive groups and cleavage of the linkers to generate glutathione on the DNA template is shown in the scheme below. It is known that glutathione binds specifically and with high affinity to Glutathione S-transferase

(GST) and is commonly used for purification of GST-fusion proteins (Amersham Pharmacia Biotech). It is also known that glutathione can be immobilized through the sulfur atom without interfering with the binding to GST. Consequently, it is possible to enrich template-displayed glutathione among other displayed molecules in a library by performing selection against GST as the target molecule. GST can be

produced in a recombinant form as described in the literature (e.g. Jemth et al. (1997) Arch. Biochem. Biophys. 348: 247-54) or be obtained from various suppliers (e.g. Sigma, product #, G5524). Alternatively, an antibody against glutathione (e.g. Abcam, product name ab64447 or Virogen, product # 101-A) can be used as the target molecule.

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Template-mediated formation of glutathione

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A microtiter plate is coated with about 1 µg streptavidin in a TBS buffer (50 mM Tris-HCI, pH 7.5, 150 mM NaCl) overnight at 4 °C. Remove the streptavidin solution and TBS buffer (other examples of blocking agent that could be used is casein, gelatine, VHS-LC-biotin as described in the literature (e.g. Ellis et al. (1998) Biochem. J. 335; glutathione for about 10-60 min and then transfer the samples from the wells to new polyvinylpyrrolidone or dried skim milk) for about 30 min. at 37 °C. Wash the plate 277-284). Free streptavidin molecules are blocked with 1 mM biotin for 5 min. and incubating at 20 °C for about 1 hour. To remove the templated molecules not coorwith TBS buffer at least three times. Add 0.1 µg biotinylated GST to the wells and incubate about 30 min at 20 °C. Remove non-bound biolinylated GST by washing excess biotin is removed by wash with TBS buffer at least six times. Add then the dinated to the immobilized GST, wash the wells with TBS buffer at least six times. wash the wells at least six times with TBS buffer. Block the wells with 2% BSA in emplated molecule library to the wells and allow binding to immobilized GST by Elute the templated molecules bound to GST by incubating with 20 mM reduced with TBS buffer at least six times. Biotinylation of GST is performed using sulfo-

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The eluted (selected) templates are amplified using two amplifying primers (forward, 5'Biotin-GCT ACT GGC ATC GGT-3'; reverse, 5'-GTA ATT GGA GTG AGC-3') with a standard PCR protocol (e.g. 5 pmol of each primer, 0.2 mM of dNTP, 2 mM of MgCl₂, and 2.5 U of thermal stable Taq polymerase). The PCR is performed with an initial denaturation at 94 °C for 5 min, 35 cycles of denaturation at 94 °C for 30 seconds, extension at 72 °C for 30 seconds, and then a final extension at 72 °C for 10 min. The 5 biotin in the forward primer is used to remove the sense strand. This is done by incubating the PCR product with streptavidin-coated magnetic beads (Dynabeads; Dynal Blotech, Norway) and the single stranded template is purified as described by the manufacturer. The purified antisense strand is finally used as the template primer together with the extension primer as describe above to generate an enriched library of templated molecules for

The selection and amplification procedure is repeated until appropriate enrichment is obtained. Enrichment can be followed by characterization (sequencing) of recovered template sequences. The nucleotide sequence of the templates is obtained using standard sequencing protocols and a DNA sequencer (e.g. MegaBase, Amersham Pharmacia Biotech). Enrichment is obtained when the number of sequences coding for glutathione (C-A-T or T-A-C in the D-D-D region of the template primer) has increased relative to other sequences in the library after the selection proce-

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another round of selection.

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This protocol describes incorporation of three different mono-nucleotide derivatives.

However, all the mono-nucleotides (including dGTP) could be used in building libraries of templated molecules as described above. Still, this will limit the number of different nucleotide derivatives to four and thus put a boundary on the library size to 4^N (where N is the number of subunits in the templated molecule). However, one may use for example di-nucleotide derivatives as building blocks in order to increase the library size to 16^N. Incorporation of di-nucleotides by polymerase has earlier been described (WO 01/16366 A2). Library diversity may be further increased using tri-nucleotides or tetra-nucleotide incorporation.

General experimental methods.

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warm up to RT and left with stirring over night. The reaction mixture was evaporated (CH₂Cl₂/MeOH=10:1 or EtOAc/heptane=2:1). The yield was in general higher than to dryness. Crude products of solid nature was recrystallized from EtOAc/heptane. A stirred solution of the amino acid (20 mmol) in CF₃COOH (10 mL) at 0 °C was slowly added (CF₃CO)₂O (24 mmol). The reaction mixture was allowed to slowly Crude products of liquid nature was purified by flash column chromatography Method 1. General procedure for M-trifluoroacetyl protection of amino acids 2 5

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Aethod 2. General procedure for N-benzyloxycarbonyl and N-vinyloxycarbonyl protection of amino acids.

A stirred solution or slight suspension of the amino acid (7.6 mmol) in sat. NaHCO3

ormation, the reaction mixture was added H2O (90 mL) and pH was adjusted to 10 pH adjusted to 2-3 using 1 M HCI (aq.) and then extracted using Et₂O or CH₂Cl₂ (3x chloroformate or vinyloxychloroformate (8.4 mmol) in CH₃CN (10 mL). The reaction using 2 M NaOH (aq.). The reaction mixture was washed with Et₂O (3x 50 mL) and mixture was left with stirring at RT over night. When TLC indicated complete trans-10 mL) was added 2 M NaOH (aq., 3 mL) and then a solution of either benzyl-

100 mL). The combined extractions were dried (MgSO₄), filtered and evaporated to

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dryness to yield a solid product, which was used without further purification. The

yield was in general higher than 70%.

- ing at RT over night. The reaction mixture was added diethyl ether (20 mL) and pH Method 3. General procedure for N-fert-butyloxycarbonyl protection of amino acids. bath), and di-tert-butyl dicarbonate was added. Further 2 M aqueous NaOH (4 mL) A slight suspension of the amino acid (15 mmol) in H₂O (5 mL) and dioxan (5 mL) vas added. The mixture was slowly heated to RT (over 5 hours), and left with stirwas added 2M NaOH (aq, 6 mL). The mixture was cooled and stirred at 0 °C (ice 5
- racted, using diethyl ether (3x 20 mL). The combined extracts were dried (MgSO4), was adjusted (from ~10 to ~3), using 2 M HCI (aq.). The aqueous phase was exitered and evaporated to dryness to yield a white solid product, which was used vithout further purification. The yield was typically 60-75 %. 2
- Method 4. General procedure for formation of NHM esters of N-protected amino 22

A stirred solution of the N-protected amino acid (0.5 mmol) and N-hydroxymaleimide bodiimide (DIC) (0.64 mmol) and the solution allowed to slowly warm up to RT and washed with a small volume of EtOAc/heptane=2/1. The filtrate was evaporated to chromatography (EtOAc/heptane=2/1), yielding the product as a white solid in typi-(0.62 mmol) in anhydrous THF (5 mL) at 0 °C under N₂ was added diisopropylcar-Imost dryness, diluted with a minimum of CH2Cl2 and subjected to flash column eft with stirring over night. The reaction mixture was filtered and the precipitate

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cally 60-70%.

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Method 5. General procedure for formation of NHM esters from carboxylic acid

sat. NaHCO₃ (aq., 2x 25 mL) and sat. NaCl (aq., 1x 25 mL). The organic phase was lowed to slowly warm up to RT and left with stirring over night. The reaction mixture dried (MgSO4), filtered, and evaporated to dryness to yield the product as a wax or was diluted with CH₂Cl₂ (16 mL) and washed with 10% citric acid (aq., 3x 25 mL), A stirred solution of N-hydroxymalelmide (4 mmol) in CH₂Cl₂ (16 mL) at 0 °C was slowly added the carboxylic acid chloride (4 mmol). The reaction mixture was alliquid in 40-60% yield. The product was used without further needed purification.

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Method 6. General procedure for S-tritylation of mercaptanes.

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A solution of the mercaptane (20 mmol) and pyridine (40 mmol) in CH₂Cl₂ (75 mL) at (eluent: CH₂Cl₂/MeOH=10/0.5). The product was isolated as an oil or a sticky wax in The volume of the reaction mixture was reduced to a minimum and then subjected RT was added tritylchloride (22 mmol) and the reaction left with stirring over night to flash chromatography (SiO₂ pretreated with pyridine prior to column packing) some instances.

Method 7. General procedure for O-acylation of 4-hydroxybenzaldehydes.

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To a stirred solution of the hydroxybenzaldehyde (20 mmol) in dry DMF (10 mL) at 0 fitered through a pad of silica. The solvent was removed in vacuo. The yield was in *C was slowly added an acid chloride (25 mmol) in diethyl ether (20 mL). The reaclion mixture was stirred at 0 °C for 15 minutes and at rt for 1 hr. Water (20 ml) was added and the reaction mixture was extracted with ether (3x10 mL). The combined vent removed in vacuo. The crude was redissolved in dichloromethane (5 mL) and organic phases was washed with water (2x10 mL), dried over MgSO4 and the solgeneral higher than 75%.

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- mL). Tosyl chloride (2.44 mmol) was added and the reaction mixture was cooled on ce. NaOH (5.5 mmol) dissolved in water (2 mL) was added dropwise and the reacion mixture was stirred at rt o/n. The reaction mixture was extracted with diethyl The corresponding polyethyleneglycol-diol (0.8 mmol), obtained as described by Baker et al. J. Org. Chem. (1999), 64, 6870-6873, was dissolved in dry THF (10 Method 8. General procedure for formation of diaminopolyethyleneglocols. ß
- phosphine (2.8 mmol) and water (2 mL) was added to the combined organic phases ether (3x5 mL) and the combined organic phases washed with NaCl (sat., 3x3 mL) and the reaction mixture was stirred o/n. IRA-120 H* (1g) was added and the reac- nL). The solution was evaporated in vacuo affording the diamino polyethylene glyion mixtured was agitated for 1 hour. The beads were filtered off, washed with direated with NaN₃ (2.8 mmol). The reaction mixture was heated to 75 °C o/n. The chloromethane (10x3 mL) and the final compound eluted with 6M HCI (aq., 10x3 and dried over MgSO4. The crude was redissolved in dry acetonitrile (3 mL) and white solid was filtered off and extracted with acetonitrile (2x2 mL). Triphenylcol in 40-50% yield. 5 9

Example 84: Prepartion of 3-phenyl-3-tertbutoxycarbonylamino-propionic acid 2,5-dioxo-2,5-dihydro-pyrrol-1-yl ester (XVI)

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Compound XVI

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The compound was prepared in two steps from the commercially available DL-3-H-NMR (CDCI₃): 7.28-7.42 (m, 5H(ar)); 6.74 (s, 2H); 5.1-5.3 (m, 2H (NH+CH)); amino-3-phenyspropionic acid by use of method 3 followed by method 4. 3.24 (dd, 1H); 3.13 (dd, 1H); 1.46 (s, 9H).

Example 85: Preparation of 3-tertbutoxycarbonylamino-butanoic acid 2.5dioxo-2,5-dihydro-pyrrol-1-yl ester

compound XVIII

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'H-NMR (CDCI₃): 6.80 (s, 2H); 4.83 (br s, 1H(NH)), 4.05-4.15 (m, 1H); 2.8-2.95 (m, The compound was prepared in two steps from the commercially available DL-3aminobutyric acid by use of method 3 followed by method 4. 2H); 1.46 (s, 9H); 2.56 (d, 3H).

Example 86: Preparation of 3-tertbutoxycarbonylamino-propionic acid 2,5dioxo-2,5-dihydro-pyrrol-1-yl ester

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Compound XVIII

H-NMR (CDCI₃): 6.80 (s, 2H); 5.09 (br s, 1H(NH)); 3.48-3.54 (m, 2H); 2.84 (t, 2H); The compound was prepared in two steps from the commercially available betaalanine by use of method 3 followed by method 4. 1.45 (s, 9H).

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Example 87: Preparation of 3-Benzyloxycarbonylamino-3-phenyl-propionic acid 2,5-dioxo-2,5-dihydro-pyrrol-1-yl ester 3-Benzyloxycarbonylamino-3phenyl-propionic acid 2,5-dioxo-2,5-dihydro-pyrrol-1-yl ester

Compound XIX

- 'H-NMR (CDCI₃): 7.55-7.20 (m, 10H); 6.75 (s, 2H); 5.55 (br., 1H); 5.35-5.25 (m, 1H); 5 , The compound was prepared in two steps from the commercially available DL-3amino-3-phenylpropionic acid by use of method 2 followed by method 4. 5.15 (s, 2H); 3.35-3.10 (m, 2H).
- Example 88: 3-Phenyl-3-vinyloxycarbonylamino-propionic acid 2,5-dioxo-2,5dihydro-pyrrol-1-yi ester 5

Compound XX

'H-NMR (CDC₁₃): 7.45-7.30 (m, 5H); 7.20 (dd, 1H); 6.75 (s, 2H); 5.75-5.60 (br., 1H); The compound was prepared in two steps from the commercially available DL-3amino-3-phenylpropionic acid by use of method 2 followed by method 4. 5.30 (q, 1H); 4.70 (d, 1H); 4.50 (d, 1H); 3.30-3.15 (m, 2H) 钇

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Example 89: Preparation of Tritvisulfanyl-acetic acid 2,5-dioxo-2,5-dihydropyrrol-1-vl ester

Compound XXI

The compound was prepared in two steps from commercially available 2mercaptoacetic acid by use of method 6 followed by method 4.

'H-NMR (CDCl₃): 7.45-7.20 (m, 15H); 6.75 (s, 2H); 3.20 (s, 2H).
Example 90: (R)-2-(2,2,2-Trifluoro-acetylamino)-3-tritylsulfanyl-pr

Example 90: (R)-2-(2,2,2-Trifluoro-acetylamino)-3-tritylsulfanyl-propionic acid 2,5-dioxo-2,5-dihydro-pyrrol-1-vl ester (XXII)

5

Compound XXII

The compound was prepared in three steps from commercially available L-cysteine by use of method 1 followed by method 6 and method 4.

Example 91: Preparation of Acetic acid 2,5-dioxo-2,5-dihydro-pyrrol-1-yl ester

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Compound XXIII

The compound was prepared in one step from commercially available acetylchloride and N-hydroxymaleimide by use of method 5.

¹H-NMR (CDCl₃): 6.75 (s, 2H); 2.35 (s, 3H).

2

Example 92: Preparation of Propionic acid 2,5-dioxo-2,5-dihydro-pyrrol-1-ylester

Compound XXIV

10 The compound was prepared in one step from commercially available propanoylchloride and A-hydroxymaleimide by use of method 5. 'H-NMR (CDCl₃): 6.75 (s, 2H); 2.65 (q, 2H); 1.80 (t, 3H). Example 93: Preparation of Butyric acid 2,5-dioxo-2,5-dihydro-pyrrol-1-yl es-

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Compound XXV

The compound was prepared in one step from commercially available butanoylchloride and A-hydroxymaleimide by use of method 5.

'H-NMR (CDCI₃): 6.75 (s, 2H); 2.60 (t, 2H); 1.80 (sxt, 2H); 1.05 (t, 3H).

Example 94: Preparation of S-Trityl-4-mercaptobenzoic acid 2,5-dioxo-2,5dihydro-pyrrol-1-yl ester

Compound XXVI

mercaptobenzoic acid, by S-tritylation according to method 6 followed by esterifica-The compound was prepared in two steps from the commercially available 4tion according to method 4.

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H-NMR-(CDCl₃): 8.75 (d, J = 8.8 Hz, 2H), 7.45-7.20 (m, 15H), 7.05 (d, J = 8.8 Hz, 2H), 6.80 (s, 2H). Example 95: Preparation of Tetrakis(aminometyI)methane tetrahydrochlorid 6

Compound XXVII

added and the white precipitate was filtered off and washed with water (100 mL) and mol) in dry pyridine (50 mL). The mixture was stirred o/n. The crude reaction mixture MeOH (200 mL). LC-MS show pentaerythritol tetratosylate. Pentaerythritol tetratosy-Tetrakis(aminomethyl)methane tetrahydrochloride was prepared by a slightly modi-Pentaerythritol (2.01 g; 14.76 mmol) was mixed with tosyl chloride (14.07 g; 73.81 ate (4.0 g, 5.31 mmol) was dissolved in dry DMF (50 mL) and NaN₃ (3.45 g; 53.1 was transferred to water (100 mL). MeOH (200 mL) and HCl conc. (80 mL) was fied method compared to Fleischer et al. J. Org. Chem. (1971), 36, 3042-44.

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phosphine (6.95 g, 26.5 mmol) and NH₃ conc. (25 mL) were added to the THF solummol) was added. The reaction mixture was heated to 100 °C o/n. Water (100 mL) THF (300 mL) was added and the diethyl either was removed in vacuo. Triphenylwas added and the reaction mixture was extracted with diethyl ether (3x100 mL).

- mL). The aqueous phase was washed with dichloromethane (3x100 mL) and evapovacuo, redissolved in dichloromethane (500 mL) and extracted with 2M HCL (2x150 rated in vacuo. MeOH (20 mL) was added and the white solid was filtered off and tion and the reaction mixture was stirred at rt o/n. The solvents were removed in washed with MeOH (2x10 mL). Yield 1.12g (76%). ß
- 14-NMR (D2O): 3.28 (s). 9

Example 96: Preparation of Propionic acid 4-formyl-phenyl ester

Compound XXVIII

The compound was prepared according to method 7 from commercially available 4hydroxybenzaldehyde. 5

'H-NMR (CDCI₃): 10.00 (s, 1H), 7.90 (d, J = 6.7 Hz, 2H), 7.31 (d, J = 6.7 Hz, 2H), 2.65 (q, J = 7.6 Hz, 2H), 1.32 (d, J = 7.5 Hz, 3H).

Example 97: Preparation of Butanoic acid 4-formyl-phenyl ester 2

Compound XXIX

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The compound was prepared according to method 7 from commercially available 4hydroxybenzaldehyde.

14-NMR (CDC13): 9.95 (s, 1H), 7.94 (d, J = 6.7 Hz, 2H), 7.28 (d, J = 6.7 Hz, 2H), 2.55 (t, J = 7.6 Hz, 2H), 1.80 (q, J = 7.6 Hz, 2H), 1.00 (d, J = 7.5 Hz, 3H).

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Example 98: Preparation of 3,6,9,12,15,18,21,24,27,30,33undecanoxapentatriacontane-1,35-diamine

Compound XXX

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Prepared according to method 8 in 48% yield. MS-H* = 545.2 (expected MS-H* = 544.6)

Example 99: Preparation of 3,6,9,12,15,18,21,24,27,30,33,36,39-Tridecanoxahentetracontane-1,41-diamine 5

Compound XXXI

(05028) Prepared according to method 8 in 40% yield. MS-H* = 633.3 (expected MS-H* = 632.8). 2

Example 100: Design and testing of oligonucleotide linkers carrying zipper boxes.

Experiments 100-1 to 100-4 were performed in order to test the efficiency of different designs of zipper boxes. The data obtained follow immediately below, then follows a discussion of the data. 22

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Materials.

Buffer A (100 mM Hepes pH= 7,5, 1 M NaCI)

Buffer B: (100 mM NaPO, pH=6, 1 M NaCI) တ

Buffer C: (100 mM NaBorate pH=9, 1 M NaCi)

Buffer D: (100 mM NaBorate pH=10, 1 M NaCl) Buffer E: (500 mM NaPO, pH=7, 1 M NaCI)

Buffer F: (500 mM NaPO, pH=8, 1 M NaCI)

9

Annealing of DNA oligonucleotides.

Mix oligos in relevant buffer and heat at 80° C then cool to 28° C (-2°C/30 sek).

5'-Labeling with 32P.

Mix 200 pmol oligonucleotide, 2 µl 10 x phosphorylation buffer (Promega cat#4103), μ T4 Polynucleotid Kinase (Promega cat#4103), 1 μl γ-3P ATP, H₂O ad 20 μl. ncubate at 37°C , 10-30 minutes. 5

PAGE (polyacrylamide gel electrophoresis).

The samples are mixed with formamide dye 1:1 (98% formamide, 10 mM EDTA, pH 0,025 % Xylene Cyanol, 0,025% Bromphenol Blue), incubate at 80°C for 2 minutes, and run on a denaturing 10% polyacrylamide gel. Develop gel using autoradiogra-ន

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phy (Kodak, BioMax film).

Oligonucteotide Building blocks

CGACCTCTGGATTGCATCGGTCATGGCTGACTGTCCGTCGAATGTGTCCAGTTACX AH37: 5'- ZGTAACTGGACTGTAAGCTGCCTGTCAGTCGGTACTGACCTGTCGAGCATCCAGCT ജ

ZGTAACACCTGTGTAAGCTGCCTGTCAGTCGGTACTGACCTGTCGAGCATCCAGCT

ZCATTGACCTGTGTAAGCTGCCTGTCAGTCGGTACTGACCTGTCGAG-AH38: 5'- AGCTGGATGCTCGACAGGTCCCGATGCAATCCAGAGGTCG

CATCCAGCT

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AH69: 5- AGZAACACCTGTGTAAGCTGCCTGTCAGTCGGTACTGACCTGTCGAG-

CATCCAGCT

AH68: 5: ZTT<u>GTAACTGGAC</u>TGTAAGCTGCCTGTCAGTCGGTACTGACCTGTCGAGCATCCAGCT

AH65: 5'. CGACCTCTGGATTGCATCGGTCATGGCTGACTGTCCGTCGAATGT<u>GTCCAGTTAC</u>TTX

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Zipper box sequences are underlined.

10 Certestrated

X= Carboxy-dT cat.no, 10-1035-

dering modifier C2.

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Y= Amino-Modifler C2 dT 10-1037-

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Z= Amino-Modifier C6 dT 10-

Experiment 100-1 (figure 56):

Mix 2 μl buffer B, 5 μl Ah36 (0,4 pmol/ul), 1 μl Ah37 (2 pmol/ul), 1 μl Ah38 (2 pmol/ul), 1 μl H₂O.

Mix 2 µl buffer B, 5 µl Ah36 (0,4 pmol/ul), 1 µl Ah37 (2 pmol/ul), 2 µl H₂O. Anneal by heating to 80° C, then cool to 44° C (-2°C/30 sek).

Add 1 µl 100 mM NHS and 1 µl 1 M EDC. Incubate at indicated temperatures (see below) for 45 minutes, then add 2 µl Buffer D. Incubate for about 2 h, and then analyze by 10% urea polyacy/amide gel electrophoresis.

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Incubation temperatures:

45 °C, 48,2 °C, 53,0 °C, 58,5 °C, 63,1 °C, 65,6 °C

Experiment 100-2 (Figure 57, A and B):

Mix 2 μl buffer B, 1 μl Ah36 (2 pmol/ul), 1 μl Ah51 (2 pmol/ul), 1 μl Ah38(2 pmol/ul),
 5 μl H₂O.

Mix 2 µl buffer B, 1 µl Ah36 (2 pmol/ul), 1 µl Ah51 (2 pmol/ul), 6 µl H₂O

Anneal by heating to 80°C, then cool to 35°C (-2°C/30 sek)(For temperatures 1 to 6), or heat to 80°C, then cool to 15°C (-2°C/30sek)(For temperatures 7 to 12).

10 Add 1 µl 100 mM NHS and 1 µl 1 M EDC. Incubate at indicated temperatures (see below) for 1 h, then add 2 µl Buffer D. Incubate for 1 h, , and then analyze by 10% urea polyacrylamide gel electrophoresis, as described above.

Incubation temperatures:

15 1) 34,9°C, 2) 36,3°C, 3) 40,3°C, 4) 45,7°C, 5) 51,0°C, 6) 55,77, 7) 14,9°C, 8) 17,8°C, 9) 22,7°C, 10) 28,3°C, 11) 31,0°C, 12) 36°C

Mix 2 µl buffer B, 0,5 µl Ah36 (2 pmol/ul), 1 µl Ah51 (2 pmol/ul), 1 µl Ah38(2 pmol/ul), 5,5 µl H₂O

20 Mix 2 µl buffer B, 0,5 µl Ah36 (2 pmol/ul), 1 µl Ah51 (2 pmol/ul), 6,5 µl H₂O
Anneal by heat at 80° C then cool to 5° C (-2°C/30 sek).
Add 1 µl 100 mM NHS and 1 µl 1 M EDC. Incubate at different temperatures (see

below) for 1 h, then add 2 µl Buffer D. Incubate for 1 h, , and then analyze by 10%

25

urea polyacrylamide gel electrophoresis.

Incubationtemperatures:

1) 5,9°C, 2) 9,9°C, 3) 12,6°C, 4) 18,3°C, 5) 23,3°C, 6) 27,9°C 7) 35,6°C, 8) 45,9°C

30 Experiment 100-3 (figure 58, A and B).

Mix 2 μ l buffer A, 1 μ l relevant oligo 1 (2 μ mol/ul), 1 μ l relevant oligo 2 (10 μ mol/ul), 1 μ l relevant oligo 3 (10 μ mol/ul), 5 μ l μ 2O. (See table below). Anneal as described

35°C for 1 sec. - repeat 99 times. Analyze by 10% urea polyacrylamide gel electro-7,7°C, 2) 15,4°C, 3) 21,0°C 4) 26,2°C for about 2 h, and 5) 10°C for 1 sec. , then Add 1 µl 100 mM NHS and 1 µl 1 M EDC. Incubate at different temperatures 1) phoresis.

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Oligo 3	Ah38	None	Ah38	None	Ah38	None	Ah38	None
Oligo 2	None	None	Ah51	Ah51	Ah67	Ah67	Ah69	Ah69
Oligo 1 (³² P)	Ah36							
Experiment	100-3-1	100-3-2	100-3-3	100-3-4	100-3-5	100-3-8	100-3-7	100-3-8

Experiment 100-4 (Figure 59).

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pmol/ul), 1 µl relevant oligo 3 (10 pmol/ul), 4,5 µl H₂O. (See table below). Anneal by heating to 80°C and then cool to 30°C or 55°C. Add 1 µl 100 mM NHS and 1 µl 1 M EDC. Incubate at 30°C or 55°C. Then analyze by 10% urea polyacrylamide gel elec-Mix 2,5 µl buffer A, 1 µl relevant oligo 1 (2 pmol/ul), 1 µl relevant oligo 2 (10 trophoresis.

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Experiment	Oligo 1 (²² P-	Oligo 2	Oligo 3
	(labelled)		
100-4-1	Ah36	Ah37	Ah38
100-4-2	Ah36	Ah37	None
100-4-3	Ah65	Ah66	Ah38
100-4-4	Ah65	Ah66	None
100-4-5	Ah36	Ah66	Ah38
100-4-6	Ah36	Ah66	None
100-4-7	Ah65	Ah37	Ah38
100-4-8	Ah65	Ah37	None

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Discussion of the results

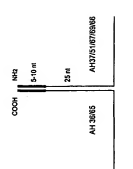
The cross-linking efficency using oligos carrying reactive groups (amine or carboxylic acid) where the linker connecting the reactive group and the annealing region

was approximately 25 nucleotides, was examined. S

rying an amine) were used. The template used (Ah38) anneals the two oligonucleo-In an experiment oligonucleotides Ah36 (carrying a carboxylic acid) and Ah67 (cartides immediately adjacent, i.e. with a spacing of zero base pairs. Under the conditions of the experiment, less than 5% cross-linking efficiency is observed, and only at the highest tested temperature (figure 58, A and B, lanes 5). 9

In order to improve the cross-linking efficiency, we introduced a so-called zipper box quences, and thus may bring the reactive groups of the two oligos into closer proxtermini that carries the reactive groups. The zipper-boxes are complementary seimity. Two different lengths of zipper boxes were tested, namely a 10'mer zipper box (Ah37/Ah36 forming a DNA duplex of 10 base pairs) and a 5'mer zipper box sequence at the 5'- and 3' end of oligos Ah67 and Ah36, respectively, the same (Ah36/51 forming a DNA duplex of 5 base pairs). See figure below.

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reactive group is attached immediately adjacent to the zipper box (Ah36, Ah37 and Ah36/Ah51), or placed two nucleotides upstream from the zipper box (Ah65/Ah66), Moreover, different designs of zipper boxes were tested, e.g. oligos in which the or placed in the middle of the zipper box (Ah67). 25

AH 38

We first tested the effect of the 5'mer zipper box on cross-linking efficiency. As can be seen, the 5'mer zipper box improves the cross-linking efficiency dramatically (figure 58, A and B, compare lanes 3 and lanes 5). Note that the template is absolutely required for cross-linking at all temperatures tested. The highest cross-linking efficiency is obtained when the temperature is cycled 99 times up and down between 10°C and 35°C (figure 58B). A high efficiency is also obtained when the temperature is kept constant at 21°C or 26°C (figure 58A and B, lanes 3). The cross-linking efficiency does not improve further at temperatures above 26°C (figure 57, A and

z,

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We next tested the efficiency of cross-linking in the 10'mer zlpper box format. Oilgos Ah36 and Ah37 were annealed to template Ah38, and the cross-linking efficiency examined at various temperatures. A surprisingly high degree of cross-linking in the absence of template was observed (figure 55, 45°C and 48.2°C). However, at temperatures above 58.5°C, no cross-linking is observed in the absence of template.

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Next, the different locations of the reactive groups relative to the zipper box was tested. As shown in figure 58, A and B, lanes 7, the cross-linking efficiency decreases dramatically when one of the two reactive groups is located in the middle of the zipper box (i.e., the reactive group is attached to a nucleotide involved in DNA double helix formation; Ah67).

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The location of the reactive groups relative to the zipper box was also tested in the context of the 10'mer zipper box. In this context, when both reactive groups are separated from the zipper box by two nucleotides (Ah65, Ah66), the efficiency of cross-linking is slightly decreased (figure 59, compare lanes 1 and 3). The cross-linking efficiency is not changed dramatically when different combinations of Ah65, Ah66, Ah36 and Ah37 are tested (i.e., when the reactive groups are placed immediately next to the zipper box, or two nucleotides upstream). Note that the template is not absolutely required at all temperatures in the context of the 10'mer zipper box. This template-independency is particularly pronounced at lower temperature (e.g.,

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Examples 101 to 104: General methods for preparation of oligonucleotide building blocks

Example 101; Procedure for transforming oligonucleotide comprising a car-

5 boxylic acid to an amino or aminomethyl terminated linker

The following oligos containing a modified nucleobase, with a carboxylic acid molety, were synthesised using the conventional phosphoramidite approach:

10 A: 5'-GCT ACT GGC XTC GGT

B: 5'-TCA CTX GCA GAC AGC

C: 5'-CGA CCT CTG GAT TGC ATC GGT CAT GGC TGA CTG TCC GTC GAA

TGT GTC CAG TTA CX

15

D: 5'-CTG GTA ACG CGG ATC GAC CTT CAT GGC TGA CTG TCC GTC GAA TGT G<u>TC CAG TTA C</u>X 20 E: 5' ACG ACT ACG TTC AGG CAA GAT CAT GGC TGA CTG TCC GTC GAA
TGT G<u>TC CAG TTA C</u>X

X was incorporated using the commercially available carboxy-dT phosphoramidite (10-1035-90 from Glen research). The underlined nucleobases represent the zipper

region.

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Schematic representation of the transformation



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figure 59, 30°C).

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An oligo (20 pmol) was mixed with a diamino compound (20 uL of a 0.1 M solution), sodiumphosphate buffer (15 uL of a 100 mM solution, pH=6), NHS (5 uL of a 100 mM solution) and EDC (5 uL of a freshly prepared 1 M solution). The mixture was left at 30 °C for 45 minutes and treated with sodium borate (20 uL of a 100 mM solution, pH=10) and left at 30 °C for additional 45 minutes. The oligo was purified by conventional EtOH precipitation. The products were and-labelled with ³²P and the purity analysed by PAGE. In all cases no starting oligo were detected and a new band, which migrated slower on the gel, appeared.

10 Examples of used diamino compounds: XXX, XXXI and the commercially available N,N'-dimethylethylenediamine (D15,780-5 from Sigma-Aldrich). Example 102: method for transforming a carboxylic acid containing oligonucleotide to a trisamine scaffold building block

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The following oligos containing a modified nucleobase, with a carboxylic acid molety, were synthesised using the conventional phosphoramidite approach:

F: 5'-GAC CTG TCG AGC ATC CAG CTG TCC ACA ATG X

20

G: 5'-GAC CTG TCG AGC ATC CAG CTT CAT GGG AAT TCC TCG TCC A<u>CA</u> <u>ATG</u> X H: 5'-GAC CTG TCG AGC ATC CAG CTT CAT GGG AAT TCC TCG TCC ACA

25 ATG XT

I: 5'-X<u>GI AAC TGG A</u>GG GTA AGC TCA TCC GAA TTC GGT ACT GAC CTG TCG AGC ATC CAG CT

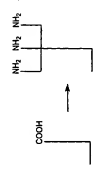
30 X was incorporated using the commercially available carboxy-dT phosphoramidite (10-1035-90 from Glen research). The underlined nucleobases represent the zipper reason.

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Schematic representation of the reaction:



An oligo containing one modified nucleobase with a carboxylic acid molety (1 nmol) was mixed with water (100 uL), hepes buffer (40 uL of a 200 mM, pH=7.5), NHS (20 uL of a 100 mM solution), EDC (20 uL of a freshly prepared 1 M solution) and the tetraamine (XXVII) (20 uL of a 100 mM solution). The reaction mixture was left on at

- 10 room temperature. The volume was reduced to 60 uL by evaporation *in vacuo*. The pure oligo was obtained by addition of NH₃ conc. (20 uL) followed by HPLC purification. It was possible to isolate a peak after approximately 6 min using the following gradient: : 0-3 minutes 100% A then 15% A and 85% B from 3-10 minutes then 100% B from 10-15 minutes then 100% A from 15-20 minutes. A = 2% acetonitrile in
- 15 10 mM TEAA and B = 80% acetonitrile in 10 mM TEAA.

After HPLC purification 2-3 pmol was end-labelled with ³²P and the purity analysed by PAGE gel (see Figure 60). The PAGE gel show the attachment of the tetraamine (XXVI) to an oligo containing a modified nucleobase with a carboxylic acid molety.

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Lane 1: Reference oligo F.

Lane 2: HPLC purified trisamine product of oligo F.

Lane 3: Reference oligo G.

Lane 4: HPLC purified trisamine product of oligo G.

25 Lane 5: Reference oligo H.

Lane 6: HPLC purified trisamine product of oligo H

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Example 103: General procedure for attachment of a functional entity to a

The following oligo containing a modified nucleobase, with a S-triphenylmethyl protected thio moiety, was synthesised using the conventional phosphoramidite ap-

proach:

J. 5: WCA TTG ACC TGT GTA AGC BTG CCT GTC AGT CGG TAC TCG ACC TCT GGA TTG CAT CGG K: 5'-WCA TTG ACC TGT CTG CCB TGT CAG TCG GTA CTG TGG TAA CGC GGA TCG ACC T 2

L: 5'-WCA TTG ACC TGA ACC ATG BTA AGC TGC CTG TCA GTC GGT ACT ACG ACT ACG TTC AGG CAA GA

5

M: 5'-WCA TTG ACC TGA ACC ATG TBA AGC TGC CTG TCA GTC GGT ACT TCA AGG ATC CAC GTG ACC AG W was incorporated using the commercially available thiol modifier phosphoramidite commercially available phosphoramidite (10-1953-95 from Glen research). The nu-(10-1926-90 from Glen research). B is an internal biotin incorporated using the cleobases which are underlined and italic indicates the zipper region.

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resuspended in TEAA buffer (200 uL of a 0.1M solution, pH=6.4). AgNO₃ (30 uL of a 1 M solution) was added and the mixture was left at room temperature for 1-2 hours. DTT (46 uL of a 1M solution) was added and left for 5-10 minutes. The reaction mix-The S-triphenylmethyl protected thio oligo (10 nmol) was evaporated in vacuo and ture was spun down (20.000 G for 20 minutes) and the supernatant was collected. The solid was extracted with additional TEAA buffer (100 ul of a 0.1 M solution, pH=6.4). The pure thio oligo was obtained by conventional EtOH-precipitation.

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Schematic representation of the reaction:

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ing the functional entity (05087) in dimethylformamide (50 ul of a 0.1 M solution) and and the loaded thio oligo was resuspended in TEAA buffer (25 uL of a 0.1M solution, left o/n at rt. The thio oligo was spun down (20.000 G for 10 minutes) and the super-The thio oligo (1 nmol) was dried in vacuo and treated with a building block compriswas spun down (20.000 G for 10 minutes). The dimethylformamide was removed natant removed. Dimethylformamide (1 mL) was added and the loaded thio oligo pH=6.4) and analysed by HPLC.

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Examples of building blocks used: XXVI, XVII, XVIII, XXIII, XXIV, XXV)

Example 104: General procedure for attachment of a functional entity to an amino or aminomethyl terminated oligo.

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The following oligo containing a modified nucleobase, with an amino group was synthesised, using the conventional phosphoramidite approach:

N: 5'-ZGT AAC ACC TGT GTA AGC TGC CTG TCA GTC GGT ACT GAC CTG

TCG AGC ATC CAG CT ଷ

commercially available amino modifier C6 dT phosphoramidite (10-1039-90 from Z contain the modified nucleobase with an aminogroup, incorporated using the Glen research)

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Furthermore, oligo C-E were transformed into the corresponding aminomethyl terminated oligo, as described earlier. The oligos were used in the following experiment represented schematically below

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An amino or aminomethyl oligo (3 pmol) was mixed with a phosphate buffer (3 uL of a 0.1 M solution, pH=6) and NaBH₃CN (3 uL of a 1 M solution in MeOH). A building block comprising the functional entity (3 uL of a 1 M solution in MeOH) was added and the mixture was left o/n at room temperature. The product formation was analysed by PAGE gel (see Figure 61).

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Examples of building blocks used: XXVIII, XXIX, and the commercially available 4-acetoxybenzaldehyde (24,260-8 from Sigma-Aldrich).

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Figure 60 shows a PAGE analysis of the loading of an oligo, containing a modified nucleobase with an amino group (comp. XXIV).

Lane 1 show the reference amino oligo (N).

Lane 2 show the amino oligo (N) after loading with a building block comprising the functional entity.

5

Lane 3 show removal of the functional entity, attached in lane 2, by treatment with pH=11 for 1 hour.

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Example 105: General procedure for the templated synthesis of an organic compound, where the scaffold and the substituent are encoded by the template:

FE NH₂ NH-FE Template

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The template oligo (1 nmol) was mixed with an thio oligo (L or M) loaded with a functional entity (XXIII or XVII, respectively, 1 nmol) and amino oligo O in hepes-buffer (20 uL of a 100 mM HEPES and 1 M NaCl solution, pH=7.5) and water (added to a final volume of 100 uL). The oligos were annealed to the template by heating to 50

- 5 °C and cooled (-2 °C/ 30 second) to 30 °C. The mixture was then left oin at a fluctuating temperature (10 °C for 1 second then 35 °C for 1 second). The oligo complex was attached to streptavidine by addition of streptavidine beads (100 uL, prewashed with 2x1 mL 100 mM hepes buffer and 1M NaCl , pH=7.5). The beads were washed with hepes buffer (1mL). The amino oligo was separated from the streptavidine
- 10 bound complex by addition of water (200 uL) followed by heating to 70 °C for 1 minute. The water was transferred and evaporated in vacuo, resuspended in TEAA buffer (45 uL of a 0.1 M solution) and product formation analysed by HPLC (see Figure 62).
- 15 Figure 62 shows the transfer of a functional entity to an oligo containing a modified nucleobase with an amino group.
- A) The top chromatogram show the reference amino oligo O: 5'-GAC CTG TCG AGC ATC CAG CTT CAT GGC TGA GTC CAC AAT GZ. 2 contain the modified nucleohase with an aminoroun incomorated using the commercially available.
 - 20 nucleobase with an aminogroup, incorporated using the commercially available amino modifier C6 dT phosphoramidite (10-1039-90 from Glen research).
- B) The middle chromatogram show the streptavidine purified amino oligo O after partial transfer of a functional entity (XXIII).

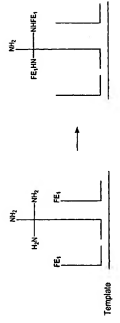
- C) The bottom chromatogram show the streptavidine purified amino oligo O after the complete transfer of a more lipophilic functional entity (XVII). The following gradient was used: 0-3 minutes 100% A then 15% A and 85% B from 3-10 minutes.
- 30 The experiment where the template oligo was omitted showed no non-templated product formation. The results indicate that the efficiency of the templated synthesis was 80-100%. The reason for less than 100% efficiency was probably due to hydrolytic cleavage of the functional entity.

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Example 106: General procedure for the templated synthesis of a scaffolded molecule, where the scaffold and two identical substituents are encoded by the template

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The template oligo (1 nmol) was mixed with two thio oligos (K and L) loaded with the same functional entity (XXVI; 1 nmol) and the trisamine oligo H (1 nmol) in hepesbuffer (20 uL of a 100 mM hepes and 1 M NaCl solution, pH=7.5) and water (added to a final volume of 100 uL). The oligos were annealed to the template by heating to 50 °C and cooled (-2 °C/30 second) to 30 °C. The mixture was then left o/n at a fluctuating temperature (10 °C for 1 second then 35 °C for 1 second). The oligo complex was attached to straptavidine by addition of streptavidine beads (100 uL, prewashed with 2x1 mL 100 mM hepes buffer and 1M NaCl, pH=7.5). The beads were washed with hepes buffer (1mL). The trisamine scaffold oligo H was separated from the straptavidine bound complex by addition of water (200 uL) followed by heating to 70 °C. The water was transferred and evaporated in vacuo, resuspended in TEAA buffer (45 uL of a 0.1 M solution) and product formation analysed by HPLC (see Figure 63).

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The HPLC chromatogram shows the transfer of two functional entities to a scaffold oligo with three amino groups.

A) The top chromatogram shows the reference scaffold oligo G.

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B) The bottom chromatogram show the streptavidine purified scaffold oligo G after the partial transfer of one (peak at 7.94 minutes) and two (peak at 10.76 minutes) identical functional entities (XXVI). The following gradient was used: 0-3 minutes 100% A then 15% A and 85% B from 3-10 minutes then 100% B from 10-15 minutes. A = 2% acetonitrile in 10 mM TEAA and B = 80% acetonitrile in 10 mM TEAA.

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Due to the lipophilic nature of the functional entities a longer retention time, in the HPLC chromatogram, of the scaffolded molecule with two functional entities compared to one functional entity, was observed. The efficency of the templated synthesis of a scaffolded molecule with the two identical functional entities (XXVI) was

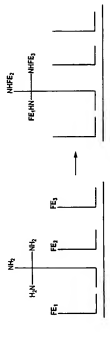
about 25% (peak at 10.76 minutes in Figure 63).

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Example 107: Procedure for the templated synthesis of a scaffolded molecule, where the scaffold and the three substituents are encoded by the tem-

plate

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- Procedure A (5-mer zipper box): The template oligo (1 nmol) was mixed with three thio oligos (J-L) loaded with three different functional entity (XVI, XVII and XVIII, respectively; 1 nmol) and the trisamine scaffold oligo H (1 nmol) in hepes-buffer (20 uL of a 100 mM hepes and 1 M NaCl solution, pH=7.5) and water (added to a final volume of 100 uL). The oligos were annealed to the template by heating to 50 °C and cooled (-2 °C/ 30 second) to 30 °C. The mixture was then left o/n at a fluctuat-
- and cooled (-2 °C/30 second) to 30 °C. The mixture was then left ofn at a fluctuating temperature (10 °C for 1 second then 35 °C for 1 second). The oligo complex was attached to streptavidine by addition of streptavidine beads (100 uL, prewashed with 2x1 mL 100 mM hepes buffer and 1M NaCl, pH=7.5). The beads were washed with hepes buffer (1mL). The trisamine scaffold oligo was separated from the strep-

(45 uL of a 0.1 M solution) and formation of the encoded molecule was identified by tavidine bound complex by addition of water (200 uL) followed by heating to 70 °C. The water was transferred and evaporated in vacuo, resuspended in TEAA buffer HPLC.

and 4-acetoxybenzaldehyde, respectively; 20 pmol) and a P^{32} end labelled trisamine Procedure B (9-mer zipper box): The template oligo (15 pmol) was mixed with three scaffold oligo I (15 pmol) in hepes-buffer (6.5 uL of a 100 mM hepes and 1 M NaCI methylamino oligos (C-E) loaded with three different functional entity (XXVII, XXIX solution, pH=7.5). The mixture was heated to 58.5 °C and left at 58.5 °C for 5 days. Formation of the encoded molecule was identified by PAGE.

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Example 108 (Model): Description of the preparation of a 3-mer \(\beta\)-amino acid

A) Synthesis of the β -amino acid building blocks

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was used as a photo cleavable N-protecting group and introduced on a β-amino acid N-terminal protection: The Nvoc group¹ (3,6-dimethoxy-6-nitrobenzyloxycarbonyl) according to the following method:

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aqueous phase was adjusted to pH = 4 with 2 M HCI (aq.) and extracted with diethyl room temperature o/n. Water (30 mL) was added and the mixture was filtered. The 3-Amino-butyric acid (147 mg, 1.43 mmol) was mixed with water (10 mL), dioxane Nvoc-Ci (1.58 mmol). 2 M NaOH was added in small portions (8 x 1.25 mL) during (10 mL) and 2 M NaOH (10 mL). The mixture was cooled to 0 °C and treated with 75 minutes. The cooling bath was removed and the reaction mixture was left at

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ether (3 x 50 mL). The solid was dissolved in water (50 mL) and diethyl ether (50 ml.). The combined organic phases were dried over MgSO, and evaporated in vacuo affording 176 mg (36 %) pure 3-(4,5-dimethoxy-2-nitro-

benzyloxycarbonylamino)-butyric acid. 1H-NMR (CDCIs): 7.72 (s, 1H), 7.02 (s, 1H), 5.51 (s, 2H), 5.40-5.30 (br s, 1H), 4.15 (m, 1H), 3.96 (s, 3H), 3.92 (s, 3H), 2.60 (d, 2H), 1.31 (d, 3H). 3-Alanin, cis-2-amino-1-cyclohexanecarboxylic acid, trans-2-Amino-1-cyclohexane carboxylic acid, cis-2-amino-1-cyclopentanecarboxylic acid, cis-2-amino-4-

- cyclohexene-1-carboxylic acid, trans-2-amino-4-cyclohexene-1-carboxylic acid, 3sminoisobutyric acid monohydrate, 3-amino-3-phenylpropionic acid, 2-fluoro-3amino-4,4,4-trifluoro butyric acid, 3-amino-4-methylpentanoic acid, DL-3aminopropionic acid hydrochloride are protected similarly. 9
- C-terminal activation: The NHM (N-hydroxymaleimide) ester of the N-Nvoc protected B-amino acid was used and prepared according to the following method, exemplified using 3-(4,5-dimethoxy-2-nitro-benzyloxycarbonylamino)-3-phenyl-propionic acid: 5

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3-(4,5-dimethoxy-2-nitro-benzyloxycarbonylamino)-butyric acid (418 mg, 1.22 mmol) was dissolved in THF (10 mL), N-hydroxymaleimide (1.22 mmol) was added and the 2H), 5.30-5.20 (br s, 1H), 4.25 (m, 1H), 3.98 (s, 3H), 3.97 (s, 3H), 2.86 (m, 2H), 1.39 (4,5-dimethoxy-2-nitro-benzyloxycarbonylamino)-butyric acid 2,5-dioxo-2,5-dihydropyrrol-1-yl ester. ¹H-NMR (CDCl₃): 7.73 (s, 1H), 7.03 (s, 1H), 6.81 (s, 2H), 5.55 (dd, mixture was cooled to 0 °C. Dicyclohexylcarbodiimide (1.22 mmol) was added and the reaction mixture was left o/n at room temperature. The solvent was removed by EIOAc-heptane (1:4 then 1:2 then 1:1) as eluent. Yield 219 mg (42 %) of pure 3evaporation in vacuo and the product isolated by silica column purification using

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¹ Burgess et al. J. Org. Chem. (1997), 62, 5165-68, Alvarez et al. J. Org. Chem. (1999), 64, 6319-28 and Pedersen et al. Proc. Natl. Acad. Sci. (1998), 95, 10523-28

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The N-Nvoc protected analogues of β-Alanin, cis-2-amino-1-cyclohexanecarboxylic acid, trans-2-Amino-1-cyclohexanecarboxylic acid, cis-2-amino-1-cyclopentanecarboxylic acid, cis-2-amino-4-cyclohexane-1-carboxylic acid, trans-2-amino-4-cyclohexane-1-carboxylic acid, 3-amino-4-cyclohexane-1-carboxylic acid, 3-amino-4-cyclohex

5 4-methylpentanoic acid, DL-3-aminoisobutyric acid monohydrate, 3-amino-3-phenylpropionic acid, 2-fluoro-3-aminopropionic acid hydrochloride, are activated

10 B) Preparation of building block oligos:

A thio oligo (1 nmol) is treated with 3-(4,5-dimethoxy-2-nitro-

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benzyloxycarbonylamino)-3-phenyl-propionic acid 2,5-dioxo-2,5-dihydro-pyrrol-1-yl ester (50 uL of 0.1 M solution in DMF). The mixture is left ofn at room temperature. The building block oligo is spinned down (20.000 G for 15 minutes) and the DMF is removed. DMF (1 mL) is added, the building block oligo is spinned down (20.000 G for 15 minutes) and the DMF is removed.

The N-Nvoc protected C-terminal NHM activated analogues of β-Alanin, cis-2-amino-1-cyclohexanecarboxylic acid, cis-2-amino-1-cyclohexanecarboxylic acid, cis-2-amino-1-cyclohexene-1-carboxylic acid, cis-2-amino-4-cyclohexene-1-carboxylic acid, 1-amino-4-cyclohexene-1-carboxylic acid, 1-amino-4-y-1-cyclohexene-1-carboxylic acid, 3-amino-4-y-1-cyclohexene-1-carboxylic acid, 3-amino-4-methylpentanoic acid, DL-3-aminoisobutyric acid monohydrate, DL-beta-aminobutyric acid, 2-fluoro-3-aminopropionic acid hydrochloride, are loaded on 11 different thio oligos similarly.

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In the following any four of the prepared building block oligos are selected and used for library production.

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C) Production of a 64-member (43) 3-mer β-peptide library:

Design of building block oligo:

Design of library setup:



15-mer Complementing element Template with three coding regions spaced by a AATAA sequence

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The sequence for the building block oligos are shown below. The nucleotides in bold constitute the complementing element and underlined the 5-mer zipperbox. ${\sf FE}^{14}$ is

the attached functional entities (4 different M-Nvoc protected β -amino acids) and B is an internal biotin.

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1) 5'-FE'-CAT IGT TIT TIT TBT TIT TIT TTT TGC ATA CAA CTA TGT A

15 2) 5'-FE²-<u>CAT 1G</u>T TTT TTT TBT TTT TTT TGC ATA CGG CTA 1GT A

3) 5'-FE³-<u>CAI 1G</u>T TTT TTT TTT TTT TTT TGC ATA CGA CTA TGT A

4) S'-FE⁴-<u>CAI 1G</u>T TIT TTT TTT TTT TTT TGC ATA CAG CTA TGT A

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5) 3'-FE1-GTA ACT TTT TTT TTT TBT TTT TTT TAT GCG TAA AGC CAT G

6) 3'-FE²-GIA ACT TIT TIT TIT TBT TIT TIT TAT GCG TGG AGC CAT G

25 7) 3'-FE³-GTA ACT TIT TIT TIT TIT TIT TAT GCG TGA AGC CAT G

8) 3'-FE4-GIA ACT TIT TIT TIT TIT TIT TIT TAT GCG TAG AGC CAT G

mM hepes buffer and 1 M NaCI, pH = 7.5). Water is added to a final volume of 1000 ul. The oligos are annealed to the templates by heating to 50 °C and cooled (-2 °C/ 64 template oligos (2 pmol each) consisting of 3 coding regions are mixed with four lamp, pyrex filter, cutoff<300 nm) for 1-2 hours. The mixture is left o/n at a fluctuat-30 second) to 20 °C. The Nvoc-protecting groups are removed by degassing thordifferent building block oligos (1-4, 200 pmol each) and hepes buffer (20 ul., 100 oughly with Ar, followed by exposure to a mercury lamp (450 W HPLC mercury ing temperature (10 °C for 1 second then 35 °C for 1 second).

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lirst template (3'-CGT ATG TTG ATA CAT AAT AAC GTA TGT TGA TAC ATA ATA separate experiments, where only one template is used in each experiment. The Formation of the encoded molecules in the library production is addressed in two ACG TAT GTT GAT ACA T) encode for the formation the 3-mer β-peptide of βalanin and the other template (3-CGT ATG CCG ATA CAT AAT AAC GTA TGC CGA TAC ATA ATA ACG TAT GCC GAT ACA T) for the formation of 3-mer \(\beta \)peptide of 3-amino-4,4,4-trifluorobutyric acid.

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A template oligo (2 pmol) consisting of 3 coding regions are mixed with four different with Ar, followed by exposure to a mercury lamp (450 W HPLC mercury lamp, pyrex buffer (1mL). The building block oligo, containing the encoded product, is separated oligos are annealed to the templates by heating to 50 °C and cooled (-2 °C/ 30 seclure (10 °C for 1 second then 35 °C for 1 second). The oligo complex is attached to building block oligos (1-4, 200 pmol each) and hepes buffer (20 u.l., 100 mM hepes ilter, cutoff<300 nm) for 1-2 hours. The mixture is left o/n at a fluctuating temperaond) to 20 °C. The Nvoc-protecting groups are removed by degassing thoroughly 100 mM hepes buffer and 1M NaCI, pH=7.5). The beads are washed with hepes buffer and 1 M NaCl, pH = 7.5). Water is added to a final volume of 100 uL. The streptavidine by addition of streptavidine beads (100 ul., prewashed with 2x1 ml. from the streptavidine bound complex by addition of water (200 ul.) followed by heating to 70 °C. The water is transferred and product formation verified by MS

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Example 109 (Model): Description of the preparation of a 3-mer B-amino acid library

A) Synthesis of the β-amino acid building blocks

was used as a photo cleavable M-protecting group and introduced on a B-amino acid N-terminal protection: The Nvoc group (3,6-dimethoxy-6-nitrobenzyloxycarbonyl) according to the method described above.

Mol. Cryst. Liq. Cryst. Sci. Technol. Sect. A (1996), 280, 17-26), according to the follow-C-terminal activation: The M-Nvoc protected β-amino acid was activated using the known 1-(4-hydroxy-phenyl)-pyrrole-2,5-dione (Choi et al. 2

room temperature o/n. The solvent is evaporated in vacuo and the pure product (3mmol). The cooling bath is removed after 1 hour and the reaction mixture is left at dimethoxy-2-nitro-benzyloxycarbonylamino)-butyric acid (1mmol) is dissolved in THF (3 mL). The solution is cooled to 0 °C and treated dropwise with DIC (1.2 (4,5-dimethoxy-2-nitro benzyloxycarbonyl-amino)-butyric acid 4-(2,5-dioxo-2,5-1-(4-Hydroxy-phenyl)-pyrrole-2,5-dione (1 mmol), NHS (1.0 mmol) and 3-(4,5dihydro-pyrrol-1-yl)-phenyl ester) is isolated by silica column purification using EtOAc-heptane (1:4 then 1:2 then 1:1) as eluent.

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The M-Nvoc protected analogues of B-Alanin, cis-2-amino-1-cyclohexanecarboxylic cyclopentanecarboxylic acid, cis-2-amino-4-cyclohexene-1-carboxylic acid, trans-2acid, trans-2-Amino-1-cyclohexanecarboxylic acid, cis-2-amino-1-

amino-4-cyclohexane-1-carboxylic acid, 3-amino-4,4,4-trifluorobutyric acid, 3-amino-4-methylpentanoic acid, DL-3-aminoisobutyric acid monohydrate, DL-beta-aminobutyric acid, 2-fluoro-3-aminopropionic acid hydrochloride, are C-terminal activated similarly.

B) Preparation of building block oligos:

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A thio oligo (2 nmol) in water (25 uL) is treated with 3-(4,5-dimethoxy-2-nitrobenzyloxycarbonyl-amino)-butyric acid 4-(2,5-dioxo-2,5-dihydro-pyrrol-1-y/)-phenyl ester (25 uL of a 10 mM solution in MeOH). The mixture is left oin at room temperature. The building block oligo is purified by a conventional EtOH-precipitation. The pellet is washed with dichloromethane (3 x 300 uL) and dried *in vacuo*.

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The N-Nvoc protected C-terminal activated analogues of β-Alanin, cis-2-amino-1-cyclohexane-carboxylic acid, trans-2-Amino-1-cyclohexanecarboxylic acid, trans-2-amino-1-cyclopentane-carboxylic acid, cis-2-amino-1-cyclohexene-1-carboxylic acid, trans-2-amino-4-cyclohexene-1-carboxylic acid, 3-amino-4-cyclohexene-1-carboxylic acid, 3-amino-4-acid, acid, 2-funoro-3-aminolsobutyric acid monohydrate, DL-beta-aminobutyric acid, 2-fuoro-3-aminopropionic acid hydrochloride, are loaded on 11 different thio oligos similarly.

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Any four of the prepared building block oligos are selected and used for library production as described above.

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Example 110 (Model); In the following, a library preparation method based on oligonucleotide templates and 5-phophoimidazolid nucleoside building blocks is described.

Preparation of building blocks

Step A: Preparation of an ester linker with a terminal alkyne.

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The acid derivative (10.37 mmol) is dissolved in DCM (20 mL) and cooled to 0 °C on an ice bath. EDC (12.44 mmol, 1.2 equiv) is added followed by DMAP (1.04 mmol, 0.1 equiv) and the alcohol (15.55 mmol, 1.5 equiv) in DCM (5 mL). After 1h reaction on ice bath, the mixture is allowed to come to 20 °C and left to react 16h. Volatiles are removed and the residue is taken up in diethylether (150 mL) and HCI (aq. 0.1 M, 75 mL). The phases are separated and the organic phase is first washed with a mixture of NaHCO₃ (sat, 35 mL) and water (35 mL) then with water (75 mL). Upon evaporation of diethylether, the product is azeotropically dried using toluene (2x120 mL) affording the desired ester that may be purified by chromatography if necessary.

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Step B. Introduction of protective groups on lodo substituted nucleosides.

The nucleoside (5.65 mmol, 1eq), TBDMS-CI (2.04g, 13.56 mmol, 2.4 eq) and imidazole (1.85g, 27.11 mmol, 4.8 eq) are mixed in DMF(20 mL) and stirred at 25 °C overnight. EtOAc (400 mL) is added and the organic phase is washed with a mixture of NH₄CI(aq) (sat, 40 mL) + H₂O (40 mL) followed by H₂O (80 mL). The organic phase is stripped and the residue is taken up in toluene, filtered and stripped to leave the desired protected nucleoside. The compound may be further purified by recrystallization.

Step C. Sonogashira coupling of protected lodo substituted nucleosides and terminal alkynes

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A DMF solution (20 mL) of the protected iodo substituted nucleoside (3.4 mmol), the alkyne (6.9 mmol, 2 eq.), DIEA (2.5 mL) is purged with Ar for 5 min. Tetrakis triphenylphosphine palladium (0.3 mmol, 0.1 eq) and Cul (0.7 mmol, 0.2 eq) is added and the mixture is heated to 50 °C and kept there for 20 h. Upon cooling, the mixture is added 700 mL diethylether. The organic phase is washed with ammonium chloride (sat, aq, 250 mL) and water (250 mL). Evaporation of volatiles followed by stripping with rolluene (400 mL) affords the desired modified nucleoside that is purified by column chromatography (sitica gel, Heptane/Ethyl acetate eluent).

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Step D: Removal of OH protective groups

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A THF solution of the above product (1.8 mmol in 30 mL) is added acetic acid (0.8 mL, 14.1 mmol, 8 eq) and tetrabutylammonium fluoride (7 mmol, 4 eq). Upon stirring at 20 °C for 20 h, volatiles are removed in vacuo and the residue is purified by column chromatography (silica gel, DCM/Methanol eluent).

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Step E: Mono-phosphate synthesis

A slurry of the modified nucleoside obtained in step D (1.65 mmol) in trimethyl phosphate (5 mL) is cooled to 0 °C and added phosphoroxytrichloride (190 uL, 307 mg, 2 mmol, 1.2 eq). The reaction is kept at 0 °C for 2h. Tributyl amine (1mL) is added and the reaction is allowed to come to 20 °C. Another portion of tributyl amine (1.3 mL) is added to raise pH, followed by water. Volatiles are removed in vacuo and the residue may be purified using lon-exchange chromatography (Sephadex A25, tetraethylammonium bromide buffer 0.05-1.0 M, pH 7).

10 Step F: Phosphoimidazolid synthesis

To a solution of the above mono-phosphate derivative (0.1 M) is added 2-methylimidazole (0.5 M) and EDC (0.5 M) at pH 6.5 and 0 °C. The reaction is stirred for 2h maintaining a temperature of 0 °C. The mixture may be used directly in library synthesis. [Visscher,1988; Journal of Molecular Evolution; 3-6]

15 Alternatively, treatment of phophates with carbonyl diimidazole also affords phophoimidazolides. [Zhao; 1998; J. Org. Chem.; 7568-7572]

A collection of building blocks

ment by means of an carboxylic ester and may be synthesized as described above. In the the scheme below a number of building blocks useful for library synthesis is shown. All building blocks have functional entities attached to the recognition ele-

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Library preparation

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The scheme below exemplifies the process of making a library of polyamides using oligonucleotide templates and phosphoimidazolid building blocks shown above. An oligonucleotide primer sequence with a sequence modifier carrying an (optionally) anneated to the templates used in the library. Further, another oligonucleotide seprotected amine (e.g. Glen Research Amino-Modifier C2 dT, cat no 10-1019-) is

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template coding the building block incorporation. For clarity, the bases of the phosquence is annealed as a terminating sequence thus exposing only the part of the phoimidazolid building blocks have been replaced with large bold letter codes.

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ncorporation

Typical conditions for oligomerisation of building blocks on the template are 0.05 M templates, 0.1-0.2 M building blocks, in a 0.2 M 2,6-lutidine-HCl buffer adjusted to pH = 7.2 buffer containing 1.0 M sodium chloride 0.2 M magnesium chloride. The temperature is kept at 0 °C for 1-21 days. [Inoue;1984; Journal of Molecular Biology;669-676]. The oligonucleotide complexes may be purified using micro-spin gel filtration (BioRad).

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Amine deprotection

Cbz protection groups may be removed by a variety of methods, [Greene;1999;]

Due to its mildness, catalytic reduction is often the method of choice. Combining an insoluble hydrogenation catalyst e.g. Pd/Al₂O₃, Pd/CaCO₃, Pd/C, PtO₂, or a soluble one e.g. Wilkinsons catalyst and a hydrogen source exemplified but not limited to H₂, ammonium formiate, formic acid, 1,4-cyclohexadien, and cyclohexane in a suitable solvent like water, methanol, eithanol, dimethylformamide, dimethylsulfoxide, ethylen glycol, acetonitril, acetic acid or a mixture of these with the oligo nucleotide

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complexes removes the Cbz protective groups.

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NaCH pH 10-12

Di-amines are linked together using di-carboxylic acids, a peptide coupling reagent optionally in the presence of a peptide coupling modifier in a suitable solvent like water, methanol, ethanol, dimethylformamide, dimethylsulfoxide, ethylene glycol, acetonitrile or a mixture of these. To an aqueous buffered solution (10uL, 1M NaCl, 100-500 mM buffer pH 6-10, preferably 7-9) of oligonucleotide complexes (0.1-100 uM, preferably 0.5-10 uM) carrying free di-amines is added a di-carboxylic acid (0.1mM-100mM, preferably 1-10 mM) exemplified by but not limited to oxalic, malonic, succinic, pentanediolc- or hexanedioic acid, phthalic, isophthalic, terephthalic acid, N-protected glutamic acid or N-protected aspartic acid mixed with a peptide coupling reagent (0.1 mM – 100 mM, preferably 1-10 mM) exemplified by but not limited to EDC, DCC, DIC, HATU, HBTU, PyBoP, PyBroP or N-methyl-2-chloropyridinium tetrafluoroborate and a peptide coupling modifier (0.1 mM-100 mM, preferably 1-10 mM) exemplified by but not limited to NHS, sulpho-NHS, HOBt, HOAt, DhbtOH in a suitable solvent (1uL) e.g. water, methanol, ethanol, dimethylformamide, dimethylsulfoxide, ethylene glycol, acetonitrile or a mixture of these.

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Reactions run at temperatures between -20 °C and 100 °C, preferably between 0 °C and 60 °C. Reaction times are between 1h and 1 week, preferably 1h-24h. The above procedure exemplifies the polymerisation on a 11 uL scale, but any other reaction volume between 1.1 uL and 1.1L may be employed.

Linker cleavage

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The ester linkages are cleaved with aqueous hydroxide at pH 9-12 at room temperature, 16 h in a suitable solvent like water, methanol, ethanol, dimethylformamide, dimethylsulfoxide, ethylene glycol, acetonitrile or a mixture of these.

MS-Analysis

10 Library members may be analyzed using Mass Spectroscopy.

In the above sequence, diamines carry Cbz protection groups and are deprotected on the oligonucleotide. Other protection schemes may also be relevant for amine protection. [Greene and Wuts;1999;] In some cases it may suffice running the sequence with building blocks that do not carry protective groups on the amines, hence eliminating the amine deprotection step. The described procedure for templated library synthesis may also employ the use of modified di- and tri-nucleotides as well as modified nucleic acid analogues like morpholinos, LNA and PNA. In the latter case reaction conditions during incorporation should be changed to accommodate peptide coupling reactions. [Schmidt;1997; Nucleic Acids Research; 4792-4796] Examples of such alternative building blocks are shown in the scheme below. Synthesis of the modified PNA units compared to ordinary PNA units differs only in the use of modified bases.

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Further, instead of using 5'-phospholmidazolide-nucleosides, a mixture of bis-3',5' phospholmidazolide-nucleosides[Visscher and Schwartz;1988; Journal of Molecular Evolution; 3-6] and nucleosides may be employed in library production, see below. Alternating incorporation of each building block type is required, but due to the reversibility of the recognition step and the fact that no reaction takes place if for instance two bis-3',5' phosphoimidazolide-nucleosides are placed next to each other all that is necessary is that both building block types are present in the mixture.

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References

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- 4cids Research 1997, 25, 4792-4796.

Example 111 (Model): Synthesis of a library of templated molecules by nonenzymatic ligation of dinucleotides comprising functional entities 15

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Several systems have been developed that enable the non-enzymatic chemical ligation of nucleotides and oligonucleotides on nucleic acid or PNA templates (Xu et al., 2001, Nat Biotechnol 19, 148-152; 2000, J Am Chem Soc, 122, 9040-41). One protocol describes the autoligation of 3'-phosphothioate and a 5'moiety comprising an iodine leaving group as shown below.

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synthesised. Each di-nucleotide comprises a 5'-iodo- and a 3'-phosphothioate group ional entity was synthesised by modified phosphoamidite nucleotide chemistry as The non-enzymatic ligation protocol can be used for the templated synthesis of a described below. 4 di-nucleotide building blocks with the sequences dUdNp were library of molecules. Here, a set of dinucleotides each comprising a unique funccapable of forming a covalent bond with a neighbouring reactive group. 9 5

incorporation of di-nucleotides on a DNA template:

where X denotes deoxythymidine-C6-NH2, Glen Research Cat#: 10-1035-90) and B pmol each of extension primers A (5'-GCTACTGGCATCGXG-3'-phosphothioate,

mM HEPES-KOH, pH=7.5, 5 mM MgCl₂, 100 mM, KCl and incubated at 80 °C for 2 eaction mixture is incubated at 4 °C for 30 min followed by a brief heating to 25 °C or 30 seconds. The reaction mixture is subjected to successive temperature oscilla-5'GCTGTCTGCAAGTGCNANACACGATGCCAGTAGC-3') in a binding buffer: 50 ion cycles for 24 hours. This step promotes the chemical ligation between correctly minutes before slowly cooling down to 20 °C. The binding of primer A and B to the emplate forms a double stranded DNA complex with a central 4 nucleotide singlestranded segment as shown below. 10 pmol of di-nucleotides are added and the 5'-lodo-GCACTTGCAGACAGC-3') are annealed to a template oligo annealed dinucleotides and the primers A and B. ឧ 22

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Following template complementation and chemical ligation, dinucleotides and buffer are removed by micro-spin gelfiltration (Biorad).

The DNA complexes comprising the functional entities are incubated in a buffer 20 50°C for 15 min before addition of equimolar HCI. The sample is transferred to a (BS₃, Pierce) is added and the sample is incubated at 30 °C for 2-8 hours. Buffer molecules are activated by cleavage of the ester linkages using 0.2 M NaOH at and excess BS₃ are removed by micro-spin gelfiltration (Biorad). The templated Cross-linking of functional entities and activation of templated molecules. mM HEPES-KOH pH = 7.5, 100 mM KCI. 5 mM Bis[Sulfosuccinimidyl]suberate suitable buffer by dialysis.

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each linked to their template applicable for selection/amplification experiments. Larger libraries can be synthesised using tri-, tetra-, or other oligonucleotides comprising functional entities and/or by increasing the number of building blocks to be cou-This protocol allows for the synthesis of a small library of 16 different molecules pled by non-enzymatic ligation on the DNA template.

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Synthesis of building blocks:

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Synthesis of 5'-iodo-3'-phosphonothioate dimers with a functional entity attached

coupling to introduce 5-I-dU; d) CF₃COOH then I₂/pyridine/water; e) FE-spacer and a) Conventional phosphoramidite coupling; b) Se in pyridine; c) phosphoramidite Pd(0) in THF-Et₃N; f) Ph₃P and I₂ in DMF; g) photolysis >300 nm.

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B equals either A, T, G or C properly protected with the photolabile protecting group Nvoc². Linker equals a photolabile CPG solid support.³ R equals a photolabile phosphate protecting group.4

Alvarcz et al. J. Org. Chem. (1999), 64, 6319-28 Pirrung et al. J. Org. Chem. (1998), 63, 241-46

Examples of attachment points (indicated by an arrow) of the linker on the nucleo-

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Examples of linker (indicated by dotted ring) and functional entity

Example 112: Ligation of ONA oligonucleotides, derivatized at the central nucleo-

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Givens and Kueper Chem. Rev. (1991), 93, 55

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DNA ligase, oligo-derivatives Ah17 and Ah19 and were annealed to templates Ah18 In order to examine the substrate efficiency of various DNA oligo-derivatives for T4 appropriate oligo. The oligo-derivatives contain a modified nucleotide at the central and Ah20, respectively. Each of the templates contain two annealing sites for the nucleotide position (see figure below).

The reaction may be schematically represented as indicated below:

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NH2		
ZH2		
	Ligase	
NH2	(AH17, AH19]	AH18, AH20)
NH2	(AH17, AH19)	(AH

X= Amino-Modifier C6 dT 9

Ah 17: 5'- CACXGAA

Ah 18: 5'- TCGGATTCAGTGTTCAGTGCGTAG

Ah 19: 5'- TGCACXGAAGC

Ah20: 5'- TCGGAGCTTCAGTGCAGCTTCAGTGCACGTAG

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DNA Ligase (TAKARA, code #6022). Incubate at 4,7°C for about 48h. Then analyze abelled) (1 pmol/µl). Anneal by heating to 80°C and then cool to 10°C. Add 3 µl T4-Mix 0,5 µl buffer A, 0,5µl Ah18 or Ah20 (1 pmol/µl), and 2 µl Ah17 or Ah18 (32Pby 10% urea polyacrylamide gel electrophoresis. As seen in figure 64, the DNA ligase is able to efficiently ligate both oligo-derivatives tested, i.e. even for the shortest oligo (Ah17), with a length of 7 nucleotides, and a modification at position 4, ligation goes to approximately 50% completion.

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Claims

 A method for synthesising a templated molecule comprising a plurality of functional groups, said method comprising the steps of

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providing at least one template comprising a sequence of n coding elements,

wherein each coding element comprises at least one recognition group capable of recognising a predetermined complementing element, and

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wherein n is an integer of more than 1,

providing a plurality of building blocks, wherein each building block ≘

comprises

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recognition group capable of recognising a predetermined coding a) at least one complementing element comprising at least one element,

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 b) at least one functional entity comprising at least one functional group and at least one reactive group, and

c) at least one linker separating the at least one functional entity from

the at least one complementing element,

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contacting each of said coding elements with a complementing element capable of recognising said coding element,

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optionally, obtaining a complementing template, and 2

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obtaining a templated molecule comprising covalently linked, functional functional group of at least one functional entity to a functional group of groups by linking, by means of a reaction involving reactive groups, a another, functional entity, 5

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wherein the templated molecule is capable of being linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, and

- wherein the synthesis of the templated molecule does not involve ribosome mediated translation of a nucleic acid. ß
- 2. Method of claim 1, wherein the templated molecule is linked by means of a single linker to the complementing template or template that templated the synthesis of the templated molecule.

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- 3. The method of claim 1, wherein steps iii) through v) are repeated.
- 4. The method of claim 1, wherein the template comprising n coding elements is a linear sequence of coding elements.

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5. The method of claim 1, wherein the template comprising n coding elements is

branched.

- from 2 to 40, for example from 2 to 30, such as from 2 to 20, for example from 2 example from 2 to 100, such as from 2 to 80, for example from 2 to 60, such as as from 2 to 4, for example 2, such as from 3 to 100, for example from 3 to 80, to 15, such as from 2 to 10, such as from 2 to 8, for example from 2 to 6, such The method of claim 1, wherein n preferably has a value of from 2 to 200, for ន
- as from 4 to 15, for example from 4 to 10, such as from 4 to 8, such as from 4 to from 3 to 20, such as from 3 to 15, for example from 3 to 15, such as from 3 to example 3, such as from 4 to 100, for example from 4 to 80, such as from 4 to 60, such as from 4 to 40, for example from 4 to 30, such as from 4 to 20, such such as from 3 to 60, such as from 3 to 40, for example from 3 to 30, such as 10, such as from 3 to 8, for example from 3 to 6, such as from 3 to 4, for

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example from 5 to 6, for example 5, such as from 6 to 100, for example from 6 to from 5 to 60, such as from 5 to 40, for example from 5 to 30, such as from 5 to 80, such as from 6 to 60, such as from 6 to 40, for example from 6 to 30, such 20, for example from 5 to 15, such as from 5 to 10, such as from 5 to 8, for 33

6, for example 4, for example from 5 to 100, such as from 5 to 80, for example

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is from 6 to 20, such as from 6 to 15, for example from 6 to 10, such as from 6 from 7 to 60, such as from 7 to 40, for example from 7 to 30, such as from 7 to to 8, such as 6, for example from 7 to 100, such as from 7 to 80, for example 20, for example from 7 to 15, such as from 7 to 10, such as from 7 to 8, for

example from 10 to 100, such as from 10 to 80, for example from 10 to 60, such axample 7, for example from 8 to 100, such as from 8 to 80, for example from 8 is from 10 to 40, for example from 10 to 30, such as from 10 to 20, for example to 60, such as from 8 to 40, for example from 8 to 30, such as from 8 to 20, for example from 8 to 15, such as from 8 to 10, such as 8, for example 9, for

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example from 12 to 30, such as from 12 to 20, for example from 12 to 15, such from 10 to 15, such as from 10 to 12, such as 10, for example from 12 to 100, as from 14 to 100, such as from 14 to 80, for example from 14 to 60, such as rom 14 to 40, for example from 14 to 30, such as from 14 to 20, for example such as from 12 to 80, for example from 12 to 60, such as from 12 to 40, for

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from 14 to 16, such as from 16 to 100, such as from 16 to 80, for example from 16 to 60, such as from 16 to 40, for example from 16 to 30, such as from 16 to

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example from 20 to 100, such as from 20 to 80, for example from 20 to 60, such 20, such as from 18 to 100, such as from 18 to 80, for example from 18 to 60, such as from 18 to 40, for example from 18 to 30, such as from 18 to 20, for

22 to 40, for example from 22 to 30, such as from 22 to 25, for example from 25 as from 20 to 40, for example from 20 to 30, such as from 20 to 25, for example o 100, such as from 25 to 80, for example from 25 to 60, such as from 25 to 40, rom 22 to 100, such as from 22 to 80, for example from 22 to 60, such as from for example from 25 to 30, such as from 30 to 100, for example from 30 to 80,

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from 40 to 60, such as from 40 to 50, for example from 40 to 45, such as from 45 as from 35 to 40, for example from 40 to 100, such as from 40 to 80, for example to 100, for example from 45 to 80, such as from 45 to 60, for example from 45 to 50, such as from 50 to 100, for example from 50 to 80, such as from 50 to 60, for as from 60 to 70, for example from 70 to 100, such as from 70 to 90, for example example from 35 to 100, such as from 35 to 80, for example from 35 to 60, such example from 50 to 55, such as from 60 to 100, for example from 60 to 80, such from 70 to 80, such as from 80 to 100, for example from 80 to 90, such as from such as from 30 to 60, for example from 30 to 40, such as from 30 to 35, for

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7. The method of claim 1, wherein the template is attached to a solid or semi-solid

8. The method of claim 1, wherein the template comprises or essentially consists of

nucleotides selected from the group consisting of deoxyribonucleic acids (DNA), LNA), and morpholinos sequences, including any analog or derivative thereof. ribonucleic acids (RNA), peptide nucleic acids (PNA), locked nucleic acids

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9. The method of claim 1, wherein the template comprises or essentially consists of

morpholinos sequence, including any analog or derivative thereof, and wherein selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and the complementing element comprises or essentially consists of nucleotides sequence, including any analog or derivative thereof. 9

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The method of claim 1, wherein the template is amplifyable.

11. The method of claim 1, wherein the template comprises a single strand of coding elements.

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The method of claim 11, wherein the single strand of coding elements is capable of forming a double helix by hybridization to a complementing template comprising a single strand of complementing elements

13. The method of any of claims 1 to 12, wherein the template comprises a priming 22

coding elements and is capable of binding a primer for initiating the incorporation 14. The method according to claim 13, wherein the priming site is upstream of the of building blocks and/or for amplification of the coding elements of the

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15. The method of claim 13, wherein the priming site is downstream of the coding

elements and is capable of binding an oligonucleotide.

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 The method of claim 15, wherein the oligonucleotide comprises an anchorage point, which can bind the templated molecule.

 The method according to claim 15, wherein the oligonucleotide is capable of stopping further incorporation of building blocks.

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- $_{\parallel}$ 18. The method of claim 15, wherein the priming site is capable of binding a reverse
- 10 19. The method of claim 1, wherein each coding element is linked to a neighbouring coding element by a covalent chemical bond.
- 20. The method of daim 1, wherein each coding element is linked to each neighbouring coding element by a covalent chemical bond.

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- 21. The method of any of claims 14 and 15, wherein said covalent chemical bond is selected from the group of covalent bonds consisting of phosphodiester bonds, phosphorothicate bonds, and peptide bonds.
- 20 22. The method of any of claims 14 and 15, wherein said covalent chemical bond is selected from the group of covalent bonds consisting of phosphodiester bonds and phosphorothioate bonds.
- 23. The method of daim 1, wherein at least one coding element is attached to a solid or semi-solid support.
- 24. The method of claim 1, wherein the coding elements are selected from the group consisting of nucleotides, including any analog or derivative thereof, amino acids, antibodies, and antigens.

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25. The method of claim 19, wherein the coding elements are selected from the group consisting of nucleotides, nucleotide derivatives, and nucleotide analogs, including any combination thereof.

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26. The method of claim 20, wherein the coding elements are selected from the group consisting of nucleotides.

- 27. The method of claim 21, wherein the nucleotides are deoxyribonucleic acids comorising a base selected from adenine (1) thumine (1) and
- comprising a base selected from adenine (A), thymine (T), guanine (G), and cytosine (C).

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28. The method of claim 21, wherein the nucleotides are ribonucteic acids comprising a base selected from adenine (A), uracil (U), guanine (G), and cytosine (C).

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- 29. The method of claim 21, wherein each nucleotide is linked to a neighbouring nucleotide by means of a covalent bond.
- 15 30. The method of daim 21, wherein each nucleotide is linked to each neighbouring nucleotide by means of a covalent bond.
- The method of any of claims 24 and 25, wherein said covalent bond is a phosphodiester bond or a phosphorothioate bond.

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- 32. The method of claim 21, wherein the coding elements are natural and non-natural nucleotides selected from the group consisting of deoxyribonucleic acids.
- 33. The method of claim 21, wherein the coding elements are natural and nonnatural nucleotides selected from the group consisting of ribonucleic acids.

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- 34. The method of claim 19, wherein the coding elements are selected from the group consisting of nucleotides, nucleotide derivatives and nucleotide analogs in which one or more of the base moiety and/or the phosphate moiety and/or the ribose or deoxyribose moiety have been substituted by an alternative molecular
- 35. The method of claim 29, wherein the complementing elements capable of interacting with said coding elements comprise or essentially consist of

entity.

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nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof

36. The method of any of claims 27 to 30, wherein each nucleotide is linked to a neighbouring nucleotide by a covalent chemical bond.

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- 37. The method of any of claims 27 to 30, wherein each nucleotide is linked to each neighbouring nucleotide by a covalent chemical bond.
- 38. The method of any of claims 31 and 32, wherein said covalent chemical bond is selected from the group of covalent bonds consisting of phosphodiester bonds and peptide bonds.

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- subunits, such as from 1 to 14 subunits, for example from 1 to 12 subunits, such essentially consists of from 1 to 100 subunits, such as from 1 to 80 subunits, for example from 1 to 5 subunits, such as from 1 to 4 subunits, for example from 1 example from 1 to 60 subunits, such as from 1 to 40 subunits, for example from as from 1 to 10 subunits, for example from 1 to 9 subunits, such as from 1 to 8 subunits, for example from 1 to 7 subunits, such as from 1 to 6 subunits, for 39. The method of claim 1, wherein the coding element preferably comprises or to 20 subunits, such as from 1 to 18 subunits, for example from 1 to 16 5 ຂ
- as from 2 to 18 subunits, for example from 2 to 16 subunits, such as from 2 to 14 subunits, such as from 2 to 40 subunits, for example from 2 to 20 subunits, such subunits, such as from 3 to 14 subunits, for example from 3 to 12 subunits, such to 3 subunits, such as from 1 to 2 subunits, for example 1 subunit, such as from example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 example from 3 to 60 subunits, such as from 3 to 40 subunits, for example from as from 3 to 10 subunits, for example from 3 to 9 subunits, such as from 3 to 8 subunits, for example from 2 to 12 subunits, such as from 2 to 10 subunits, for to 7 subunits, such as from 2 to 6 subunits, for example from 2 to 5 subunits, subunits, such as from 3 to 100 subunits, such as from 3 to 80 subunits, for 2 to 100 subunits, such as from 2 to 80 subunits, for example from 2 to 60 such as from 2 to 4 subunits, for example from 2 to 3 subunits, such as 2 3 to 20 subunits, such as from 3 to 18 subunits, for example from 3 to 16

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subunits, for example from 4 to 100 subunits, such as from 4 to 80 subunits, for эхатрle from 3 to 5 subunits, such as from 3 to 4 subunits, for example 3

as from 5 to 40 subunits, for example from 5 to 20 subunits, such as from 5 to 18 subunits, such as from 5 to 80 subunits, for example from 5 to 60 subunits, such subunits, such as from 4 to 14 subunits, for example from 4 to 12 subunits, such example from 5 to 12 subunits, such as from 5 to 10 subunits, for example from example from 4 to 60 subunits, such as from 4 to 40 subunits, for example from subunits, for example from 5 to 16 subunits, such as from 5 to 14 subunits, for as from 4 to 10 subunits, for example from 4 to 9 subunits, such as from 4 to 8 to 9 subunits, such as from 5 to 8 subunits, for example from 5 to 7 subunits, example from 4 to 5 subunits, for example 4 subunits, such as from 5 to 100 such as from 5 to 6 subunits, such as 5 subunits, for example from 6 to 100 subunits, for example from 4 to 7 subunits, such as from 4 to 6 subunits, for to 20 subunits, such as from 4 to 18 subunits, for example from 4 to 16

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as from 6 to 40 subunits, for example from 6 to 20 subunits, such as from 6 to 18 subunits, such as from 6 to 80 subunits, for example from 6 to 60 subunits, such example from 6 to 12 subunits, such as from 6 to 10 subunits, for example from 3 to 9 subunits, such as from 6 to 8 subunits, for example from 6 to 7 subunits, subunits, for example from 6 to 16 subunits, such as from 6 to 14 subunits, for such as 6 subunits, such as from 7 to 100 subunits, such as from 7 to 80

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such as from 8 to 80 subunits, for example from 8 to 60 subunits, such as from 8 to 40 subunits, for example from 8 to 20 subunits, such as from 8 to 18 subunits, as from 7 to 8 subunits, such as 7 subunits, for example from 8 to 100 subunits, example from 7 to 20 subunits, such as from 7 to 18 subunits, for example from subunits, such as from 7 to 10 subunits, for example from 7 to 9 subunits, such subunits, for example from 7 to 60 subunits, such as from 7 to 40 subunits, for for example from 8 to 16 subunits, such as from 8 to 14 subunits, for example from 8 to 12 subunits, such as from 8 to 10 subunits, for example from 8 to 9 to 16 subunits, such as from 7 to 14 subunits, for example from 7 to 12

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subunits, for example 8 subunits, such as from 9 to 100 subunits, such as from 9 subunits, such as from 9 to 10 subunits, such as 9 subunits, for example from 10 to 80 subunits, for example from 9 to 60 subunits, such as from 9 to 40 subunits, from 9 to 16 subunits, such as from 9 to 14 subunits, for example from 9 to 12 for example from 9 to 20 subunits, such as from 9 to 18 subunits, for example to 100 subunits, such as from 10 to 80 subunits, for example from 10 to 60

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subunits, for example from 3 to 7 subunits, such as from 3 to 6 subunits, for

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subunits, such as from 10 to 40 subunits, for example from 10 to 20 subunits, such as from 10 to 18 subunits, for example from 10 to 16 subunits, such as from 10 to 14 subunits, for example from 10 to 12 subunits, such as 10 subunits, from 11 to 60 subunits, such as from 11 to 40 subunits, for example from 11 to such as from 11 to 100 subunits, such as from 11 to 80 subunits, for example

20 subunits, such as from 11 to 18 subunits, for example from 11 to 16 subunits, such as from 11 to 14 subunits, for example from 11 to 12 subunits, such as

rom 12 to 100 subunits, such as from 12 to 80 subunits, for example from 12 to 50 subunits, such as from 12 to 40 subunits, for example from 12 to 20 subunits,

from 12 to 14 subunits, for example from 13 to 100 subunits, such as from 13 to such as from 12 to 18 subunits, for example from 12 to 16 subunits, such as 9

30 subunits, for example from 13 to 60 subunits, such as from 13 to 40 subunits, for example from 13 to 20 subunits, such as from 13 to 18 subunits, for example from 13 to 16 subunits, such as from 13 to 14 subunits, for example from 14 to

subunits, such as from 14 to 40 subunits, for example from 14 to 20 subunits, 100 subunits, such as from 14 to 80 subunits, for example from 14 to 60

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from 15 to 100 subunits, such as from 15 to 80 subunits, for example from 15 to 50 subunits, such as from 15 to 40 subunits, for example from 15 to 20 subunits, such as from 14 to 18 subunits, for example from 14 to 16 subunits, such as

from 16 to 100 subunits, such as from 16 to 80 subunits, for example from 16 to such as from 15 to 18 subunits, for example from 15 to 16 subunits, such as

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60 subunits, such as from 16 to 40 subunits, for example from 16 to 20 subunits, from 17 to 80 subunits, for example from 17 to 60 subunits, such as from 17 to such as from 16 to 18 subunits, for example from 17 to 100 subunits, such as

40 subunits, for example from 17 to 20 subunits, such as from 17 to 18 subunits. or example from 18 to 100 subunits, such as from 18 to 80 subunits, for 22

from 18 to 20 subunits, such as from 19 to 100 subunits, such as from 19 to 80 example from 18 to 60 subunits, such as from 18 to 40 subunits, for example

subunits, for example from 19 to 60 subunits, such as from 19 to 40 subunits, for rom 20 to 100 subunits, such as from 20 to 80 subunits, for example from 20 to 30 subunits, such as from 20 to 40 subunits, for example from 20 to 30 subunits, example from 19 to 30 subunits, such as from 19 to 25 subunits, for example

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such as from 20 to 25 subunits.

 The method of claim 39, wherein each subunit comprises or essentially consists of a nucleotide, or a nucleotide analog.

41. The method of claim 40, wherein each subunit comprises or essentially consists

of a nucleotide.

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comprising a base selected from adenine (A), thymine (T), guanine (G), and 42. The method of claim 41, wherein the nucleotide is a deoxyribonucleic acid cytosine (C).

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 The method of claim 41, wherein the nucleotide is a ribonucleic acid comprising a base selected from adenine (A), uracil (U), guanine (G), and cytosine (C).

44. The method of claim 41, wherein each nucleotide is linked to a neighbouring

nucleotide, or nucleotide analog, by means of a covalent bond 5

45. The method of claim 41, wherein each nucleotide is linked to each neighbouring nucleotide, or nucleotide analog, by means of a covalent bond. 46. The method of any of claims 44 and 45, wherein said covalent bond is selected from the group consisting of phosphodiester bonds, phosphorothioate bonds, 8

and peptide bonds.

47. The method of claim 40, wherein at least some of said nucleotides are selected

from the group consisting of nucleotide derivatives. 22 48. The method of claim 47, wherein the nucleotide derivatives are selected from the group consisting of deoxyribonucleic acid derivatives and ribonucleic acid derivatives. 49. The method of claim 39, wherein the coding element subunits are selected from the group consisting of nucleotides, nucleotide derivatives and nucleotide

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and/or a ribose molety and/or a deoxyribose moiety have been substituted by an analogs in which one or more of a base moiety and/or a phosphate moiety

alternative molecular entity.

50. The method of claim 49, wherein the complementing element subunits capable of interacting with said coding element subunits comprise or essentially consist of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof.

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- neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond. 51. The method of claim 47, wherein each nucleotide derivative is linked to a
- 52. The method of claim 47, wherein each nucleotide derivative is linked to each neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond

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- 53. The method of any of claims 51 and 52, wherein said covalent chemical bond is selected from the group of covalent bonds consisting of phosphodiester bonds,
 - phosphorothioate bonds, and peptide bonds 5

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The method according to ctaim 1, wherein a complementing template is obtained by complementing a plurality of predetermined coding elements by contacting each of said coding elements with a complementing element capable of recognising said coding element.

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- 55. The method of claim 1, wherein the complementing template comprising n complementing elements is a linear sequence of coding elements.
- 56. The method of claim 1, wherein the complementing template comprising n complementing elements is branched. 22

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57. The method of any of claims 55 and 56, wherein n preferably has a value of from 2 to 200, for example from 2 to 100, such as from 2 to 80, for example from 2 to as from 3 to 10, such as from 3 to 8, for example from 3 to 6, such as from 3 to from 2 to 6, such as from 2 to 4, such as 2, such as from 3 to 100, for example from 3 to 80, such as from 3 to 60, such as from 3 to 40, for example from 3 to 30, such as from 3 to 20, such as from 3 to 15, for example from 3 to 15, such example from 2 to 15, such as from 2 to 10, such as from 2 to 8, for example 60, such as from 2 to 40, for example from 2 to 30, such as from 2 to 20, for

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from 5 to 20, for example from 5 to 15, such as from 5 to 10, such as from 5 to 8, from 7 to 20, for example from 7 to 15, such as from 7 to 10, such as from 7 to 8, as from 35 to 40, for example from 40 to 100, such as from 40 to 80, for example for example from 5 to 6, for example 5, such as from 6 to 100, for example from example from 10 to 100, such as from 10 to 80, for example from 10 to 60, such example from 20 to 100, such as from 20 to 80, for example from 20 to 60, such 22 to 40, for example from 22 to 30, such as from 22 to 25, for example from 25 to 100, such as from 25 to 80, for example from 25 to 60, such as from 25 to 40, example from 35 to 100, such as from 35 to 80, for example from 35 to 60, such as from 10 to 40, for example from 10 to 30, such as from 10 to 20, for example as from 20 to 40, for example from 20 to 30, such as from 20 to 25, for example 4, for example 3, such as from 4 to 100, for example from 4 to 80, such as from such as 7, for example from 8 to 100, such as from 8 to 80, for example from 8 from 22 to 100, such as from 22 to 80, for example from 22 to 60, such as from example from 7 to 60, such as from 7 to 40, for example from 7 to 30, such as to 60, such as from 8 to 40, for example from 8 to 30, such as from 8 to 20, for from 14 to 16, such as from 16 to 100, such as from 16 to 80, for example from example from 5 to 60, such as from 5 to 40, for example from 5 to 30, such as example from 12 to 30, such as from 12 to 20, for example from 12 to 15, such 16 to 60, such as from 16 to 40, for example from 16 to 30, such as from 16 to from 10 to 15, such as from 10 to 12, such as 10, for example from 12 to 100, such as from 6 to 20, such as from 6 to 15, for example from 6 to 10, such as 20, such as from 18 to 100, such as from 18 to 80, for example from 18 to 60, for example from 25 to 30, such as from 30 to 100, for example from 30 to 80, 4 to 60, such as from 4 to 40, for example from 4 to 30, such as from 4 to 20, 6 to 80, such as from 6 to 60, such as from 6 to 40, for example from 6 to 30, as from 14 to 100, such as from 14 to 80, for example from 14 to 60, such as from 14 to 40, for example from 14 to 30, such as from 14 to 20, for example such as from 4 to 15, for example from 4 to 10, such as from 4 to 8, such as such as from 18 to 40, for example from 18 to 30, such as from 18 to 20, for such as from 12 to 80, for example from 12 to 60, such as from 12 to 40, for such as from 30 to 60, for example from 30 to 40, such as from 30 to 35, for from 4 to 6, such as 4, for example from 5 to 100, such as from 5 to 80, for from 6 to 8, such as 6, for example from 7 to 100, such as from 7 to 80, for example from 8 to 15, such as from 8 to 10, for example 8, such as 9, for S

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from 40 to 60, such as from 40 to 50, for example from 40 to 45, such as from 45 to 100, for example from 45 to 80, such as from 45 to 60, for example from 45 to 80, such as from 50 to 100, for example from 50 to 80, such as from 50 to 60, for example from 50 to 55, such as from 60 to 70, for example from 60 to 70, for example from 70 to 100, such as from 70 to 90, such as from 90 to 100.

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 The method of claim 1, wherein the complementing template is attached to a solid or semi-solid support.

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59. The method of claim 1, wherein the complementing template comprises or essentially consists of nucleotides selected from the group consisting of deoxyribonucleic acids (DNA), ribonucleic acids (RNA), peptide nucleic acids (PNA), locked nucleic acids (LNA), and morpholinos sequences, including any analog or derivative thereof.

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60. The method of claim 1, wherein the complementing template comprises or essentially consists of nucleotides selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof, and wherein the corresponding coding element of the template comprises or essentially consists of nucleotides selected from the group consisting of DNA, RNA, LNA and morpholinos sequence, including any analog or derivative thereof.

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61. The method of claim 1, wherein the complementing template is amplifyable.

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62. The method of claim 1, wherein the complementing template comprises a single strand of complementing elements.

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63. The method of claim 55 and 56, wherein the single strand of complementing elements is capable of forming a double helix by hybridization to a template comprising a single strand of coding elements.

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- 64. The method of any of claims 1 to 57, wherein the complementing template comprises a priming site.
- 65. The method of claim 1, wherein each complementing element is linked to a neighbouring complementing element by a covalent chemical bond.
- 66. The method of claim 1, wherein each complementing element is linked to each neighbouring complementing element by a covalent chemical bond.
- 10 67. The method of any of claims 59 and 60, wherein said covalent chemical bond is
- selected from the group of covalent bonds consisting of phosphodiester bonds, phosphorothicate bonds, and peptide bonds.

 68. The method according to any of the claims 65 to 67, wherein the phosphodiester bond is formed by a reaction between reactive groups type t, where one reactive
- 69. The method of any of claims 59 and 60, wherein said covalent chemical bond is selected from the group of covalent bonds consisting of phosphodiester bonds and phosphorothioate bonds.

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group type I is a hydroxyl group and the other is a phosphorimidazole group.

- 70. The method of claim 1, wherein at least one complementing element is attached to a solid or semi-solid support.
- 71. The method of claim 1, wherein the complementing elements are selected from the group consisting of nucleotides, including any analog or derivative thereof, amino acids, antibodies, and antigens.
- 72. The method of claim 64, wherein the complementing elements are selected from the group consisting of nucleotides, nucleotide derivatives, and nucleotide analogs, including any combination thereof.

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73. The method according to claim 72, wherein, to increase the diversity, the nucleotide derivatives comprises natural as well as non-natural derivatives.

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74. The method of claim 73, wherein the complementing elements are selected from the group consisting of nucleotides.

75. The method of claim 74, wherein the nucleotides are deoxynbonucleic acids comprising a base selected from adenine (A), thymine (T), guanine (G), cytosine (C), 7-deaza-guanine, 7-deaza-adenine.

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76. The method of claim 74, wherein the nucleotides are ribonucleic acids comprising a base selected from adenine (A), uracil (U), guanine (G), and cytosine (C).

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- 77. The method of claim 74, wherein each nucleotide is linked to a neighbouring nucleotide, or nucleotide analog, by means of a covalent bond.
- 15 78. The method of claim 74, wherein each nucleotide is linked to each neighbouring nucleotide, or nucleotide analog, by means of a covalent bond.
- 79. The method of any of claims 77 and 78, wherein said covalent bond is a phosphodlester bond or a phosphorothicate bond.

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- 80. The method of claim 72, wherein the complementing elements are natural or non-natural nucleotides selected from the group consisting of deoxyribonucleic acids.
- 25 81. The method of claim 72, wherein the complementing elements are natural or non-natural nucleotides selected from the group consisting of ribonucleic acids.
- 82. The method of claim 72, wherein the complementing elements are selected from the group consisting of nucleotides, nucleotide derivatives and nucleotide analogs in which one or more of a base moiety and/or a phosphate moiety and/or a ribose and/or a deoxyribose moiety has been substituted by an alternative molecular entity.

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83. The method of claim 82, wherein the coding elements capable of interacting with said complementing elements comprise or essentially consist of nucleotides

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selected from the group consisting of DNA, RNA, PNA, LNA and morpholinos sequence, including any analog or derivative thereof.

- 84. The method of any of claims 82 and 83, wherein each nucleotide is linked to a neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond.
- 85. The method of any of claims 82 and 83, wherein each nucleotide is linked to each neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond.
- 86. The method of any of claims 84 and 85, wherein said covalent chemical bond is selected from the group of covalent bonds consisting of phosphodiester bonds, phosphorothioate bonds, and peptide bonds.

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- 15 87. The method according to claim 1, wherein a coding element is non-covalently coupled to another coding element.
- The method according to claim 87, wherein the non-covalent coupling involves hydrogen bondings.

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89. The method of claim 1, wherein the complementing element preferably comprises or essentially consists of from 1 to 100 subunits, such as from 1 to 80 subunits, for example from 1 to 60 subunits, such as from 1 to 40 subunits, for example from 1 to 20 subunits, such as from 1 to 18 subunits, for example from 1 to 18 subunits, such as from 1 to 10 subunits, for example from 1 to 9 subunits, such as from 1 to 10 subunits, for example from 1 to 9 subunits, such as from 1 to 8 subunits, for example from 1 to 5 subunits, such as from 1 to 6 subunits, for example from 1 to 5 subunits, such as from 1 to 2 subunits, for example 1 subunit, such as from 2 to 100 subunits, such as from 2 to 80 subunits, for example 1

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example from 1 to 3 subunits, such as from 2 to 80 subunits, for example to subunit, such as from 2 to 80 subunits, for example from 2 to 60 subunits, such as from 2 to 40 subunits, for example from 2 to 20 subunits, such as from 2 to 18 subunits, for example from 2 to 16 subunits, such as from 2 to 14 subunits, for example from 2 to 12 subunits, such as from 2 to 10 subunits, for example from 2 to 9 subunits, such as from 2 to 8 subunits, for example from 2 to 7 subunits, for example from 2 to 8 subunits, for example from 8 from

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example from 2 to 5 subunits, such as from 2 to 4 subunits, for example from 2 to 3 subunits, such as 2 subunits, such as from 3 to 100 subunits, such as from

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as from 9 to 40 subunits, for example from 9 to 20 subunits, such as from 9 to 18 subunits, for example from 14 to 60 subunits, such as from 14 to 40 subunits, for subunits, for example from 15 to 60 subunits, such as from 15 to 40 subunits, for subunits, such as from 9 to 80 subunits, for example from 9 to 60 subunits, such or example from 10 to 60 subunits, such as from 10 to 40 subunits, for example subunits, for example from 11 to 60 subunits, such as from 11 to 40 subunits, for or example from 12 to 60 subunits, such as from 12 to 40 subunits, for example 16 subunits, such as from 10 to 14 subunits, for example from 10 to 12 subunits, from 14 to 16 subunits, such as from 15 to 100 subunits, such as from 15 to 80 from 10 to 20 subunits, such as from 10 to 18 subunits, for example from 10 to rom 11 to 16 subunits, such as from 11 to 14 subunits, for example from 11 to rom 12 to 20 subunits, such as from 12 to 18 subunits, for example from 12 to from 13 to 18 subunits, for example from 13 to 16 subunits, such as from 13 to subunits, for example from 10 to 100 subunits, such as from 10 to 80 subunits, subunits, for example from 8 to 12 subunits, such as from 8 to 10 subunits, for subunits, for example from 9 to 16 subunits, such as from 9 to 14 subunits, for 12 subunits, such as from 12 to 100 subunits, such as from 12 to 80 subunits, example from 14 to 20 subunits, such as from 14 to 18 subunits, for example example from 15 to 20 subunits, such as from 15 to 18 subunits, for example such as 10 subunits, such as from 11 to 100 subunits, such as from 11 to 80 example from 11 to 20 subunits, such as from 11 to 18 subunits, for example subunits, such as from 13 to 80 subunits, for example from 13 to 60 subunits, example from 8 to 9 subunits, for example 8 subunits, such as from 9 to 100 such as from 13 to 40 subunits, for example from 13 to 20 subunits, such as example from 9 to 12 subunits, such as from 9 to 10 subunits, such as 9 16 subunits, such as from 12 to 14 subunits, for example from 13 to 100 4 subunits, for example from 14 to 100 subunits, such as from 14 to 80 S 5 2 2 2

subunits, such as from 4 to 8 subunits, for example from 4 to 7 subunits, such as

rom 4 to 6 subunits, for example from 4 to 5 subunits, for example 4 subunits,

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example from 4 to 16 subunits, such as from 4 to 14 subunits, for example from

4 to 12 subunits, such as from 4 to 10 subunits, for example from 4 to 9

subunits, such as from 3 to 8 subunits, for example from 3 to 7 subunits, such as

subunits, for example 3 subunits, for example from 4 to 100 subunits, such as from 4 to 80 subunits, for example from 4 to 60 subunits, such as from 4 to 40 subunits, for example from 4 to 20 subunits, such as from 4 to 18 subunits, for

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from 3 to 6 subunits, for example from 3 to 5 subunits, such as from 3 to 4

example from 3 to 16 subunits, such as from 3 to 14 subunits, for example from

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3 to 12 subunits, such as from 3 to 10 subunits, for example from 3 to 9

subunits, for example from 3 to 20 subunits, such as from 3 to 18 subunits, for

3 to 80 subunits, for example from 3 to 60 subunits, such as from 3 to 40

as from 5 to 14 subunits, for example from 5 to 12 subunits, such as from 5 to 10

subunits, such as from 5 to 18 subunits, for example from 5 to 16 subunits, such

such as from 5 to 100 subunits, such as from 5 to 80 subunits, for example from

5 to 60 subunits, such as from 5 to 40 subunits, for example from 5 to 20

as from 6 to 14 subunits, for example from 6 to 12 subunits, such as from 6 to 10

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subunits, for example from 6 to 9 subunits, such as from 6 to 8 subunits, for

example from 6 to 7 subunits, such as 6 subunits, such as from 7 to 100

subunits, such as from 6 to 18 subunits, for example from 6 to 16 subunits, such

for example from 6 to 100 subunits, such as from 6 to 80 subunits, for example

rom 6 to 60 subunits, such as from 6 to 40 subunits, for example from 6 to 20

example from 5 to 7 subunits, such as from 5 to 6 subunits, such as 5 subunits,

subunits, for example from 5 to 9 subunits, such as from 5 to 8 subunits, for

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as from 7 to 40 subunits, for example from 7 to 20 subunits, such as from 7 to 18

example from 7 to 12 subunits, such as from 7 to 10 subunits, for example from

? to 9 subunits, such as from 7 to 8 subunits, such as 7 subunits, for example

subunits, for example from 7 to 16 subunits, such as from 7 to 14 subunits, for

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subunits, such as from 7 to 80 subunits, for example from 7 to 60 subunits, such

subunits, for example from 16 to 60 subunits, such as from 16 to 40 subunits, for rom 17 to 100 subunits, such as from 17 to 80 subunits, for example from 17 to 30 subunits, such as from 17 to 40 subunits, for example from 17 to 20 subunits rom 15 to 16 subunits, such as from 16 to 100 subunits, such as from 16 to 80 from 18 to 80 subunits, for example from 18 to 60 subunits, such as from 18 to example from 16 to 20 subunits, such as from 16 to 18 subunits, for example such as from 17 to 18 subunits, for example from 18 to 100 subunits, such as 40 subunits, for example from 18 to 20 subunits, such as from 19 to 100 ဓ 32

as from 8 to 18 subunits, for example from 8 to 16 subunits, such as from 8 to 14

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subunits, such as from 8 to 40 subunits, for example from 8 to 20 subunits, such

from 8 to 100 subunits, such as from 8 to 80 subunits, for example from 8 to 60

from 19 to 25 subunits, for example from 20 to 100 subunits, such as from 20 to 80 subunits, for example from 20 to 60 subunits, such as from 20 to 40 subunits, subunits, such as from 19 to 80 subunits, for example from 19 to 60 subunits, such as from 19 to 40 subunits, for example from 19 to 30 subunits, such as for example from 20 to 30 subunits, such as from 20 to 25 subunits.

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- 90. The method of claim 89, wherein each subunit comprises or essentially consists of a nucleotide, or a nucleotide analog.
- 91. The method of claim 90, wherein each subunit comprises or essentially consists of a nucleotide. 9
- comprising a base selected from adenine (A), thymine (T), guanine (G), and 92. The method of claim 91, wherein the nucleotide is a deoxyribonucleic acid

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- 93. The method of claim 91, wherein the nucleotide is a ribonucleic acid comprising a base selected from adenine (A), uracil (U), guanine (G), and cytosine (C).
- 94. The method of claim 91, wherein each nucleotide is linked to a neighbouring nucleotide, or nucleotide analog, by means of a covalent bond. 8
- 95. The method of claim 91, wherein each nucleotide is linked to each neighbouring nucleotide, or nucleotide analog, by means of a covalent bond.

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- 96. The method of claim 94, wherein said covalent bond is selected from the group consisting of phosphodiester bonds, phosphorothioate bonds, and peptide bonds.
- 97. The method of claim 90, wherein at least some of said nucleotides are selected from the group consisting of nucleotide derivatives ဓ္က
- 98. The method of claim 97, wherein the nucleotide derivatives are selected from the group consisting of deoxyribonucleic acid derivatives and ribonucleic acid

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nucleotide analogs in which one or more of a base moiety and/or a phosphate selected from the group consisting of nucleotides, nucleotide derivatives, and 99. The method of claim 90, wherein the complementing element subunits are moiety and/or a ribose moiety and/or a deoxyribose moiety has been

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- The method of claim 99, wherein the coding element subunits capable RNA, PNA, LNA and morpholinos sequence, including any analog or derivative essentially consists of nucleotides selected from the group consisting of DNA, of interacting with said complementing element subunits comprises or substitutetd by an alternative molecular entity. 9
- The method of claim 93, wherein each nucleotide derivative is linked to a neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond. ₽. 5
- The method of claim 93, wherein each nucleotide derivative is linked to each neighbouring nucleotide, or nucleotide analog, by a covalent chemical bond. ₽ 2
- chemical bond is selected from the group of covalent bonds consisting of The method of any of claims 97 and 98, wherein said covalent phosphodiester bonds, phosphorothioate bonds, and peptide bonds. 5.

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- The method of claim 1, wherein the complementing elements are selected from nucleotides, and the complementing elements are linked enzymatically. ₹ 22
- from the group consisting of template-dependent DNA- and RNA-polymerases, including reverse transcriptases, DNA-ligases and RNA-ligases, ribozymes and The method according to daim 100, wherein the enzyme is selected Sequenase, Taq DNA polymerase, Klenow Fragment (Large fragment of DNA Transcriptase, T7 RNA polymerase, T7 RNA polymerase mutant Y639F, deoxyribozymes, including HIV-1 Reverse Transcriptase, AMV Reverse ജ

polymerase I), DNA-ligase, T7 DNA polymerase, T4 DNA polymerase, T4 DNA

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Ligase, E. coli RNA polymerase, rTh DNA polymerase, Vent DNA polymerase, Ptu DNA polymerase, Tte DNA polymerase, repair polymerase, and ribozymes with ligase or replicase activities.

- 106. The method of daim 105, wherein the enzyme is selected from the group consisting of HIV-1 Reverse Transcriptase, AMV Reverse Transcriptase, T7 RNA polymerase, T7 RNA polymerase mutent Y639F, Sequenase, Taq DNA polymerase, Klenow Fragment (Large fragment of DNA polymerase), DNA-ligase, T7 DNA polymerase, T4 DNA polymerase, and T4 DNA Ligase.
- 107. The method of any of claims 104 to 106, wherein the nucleotides form a complementing template upon incorporation.

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108. The method according to any of the claim 104 to 107, wherein a primer is annealed to a priming site on the template and a suitable polymerase or transcriptase extents the primer by incorporation of building blocks to obtain a complementing template.

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- The method according to claim 108, wherein the building block is a
 mono-, di- or oligonucleotide derivative.
- 110. The method according to claim 109, wherein the building block is a mononucleotide derivative.
- The method according to any of the claims 109 to 110, wherein the functional entity is attached to the nucleobase moiety through a linker.
- 112. The method according to claim 107, wherein the linker comprises a triple bond.

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113. The method according to any of the claims 108 to112, wherein the building block is designed such that the functional entity protrudes into the major groove of a nucleic acid helix, when incorporated in a complementing template.

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- 114. The method according to any of the claims 104 to 106, wherein the complementing elements are selected from oligonucleotides and the complementing elements are linked enzymatically using a ligase.
- 5 115. The method according to claim 114, wherein the complementing element oligonucleotides each comprise three or more nucleic acid monomers.
- 116. The method according to claim 115, wherein the complementing element is an oligonucleotide comprising 4 to 100 nucleic acid monomers.

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- 117. The method according to claim 1, wherein a complementing element anneals to the coding element, without being coupled to another complementing element.
- 15 The method according to claim 117, wherein the complementing element comprises an oligonucleotide of 6 to 100 nucleotide monomers.
- 119. The method according to claim 119, wherein the oligonucleotide comprises 10 to 50 nucleotide monomers.

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- 120. The method according to any of the claims 114 to 119, wherein the langth is at least half the length of the complementing element.
- 121. The method according to any of the claims 114 to 120, wherein the
 - 25 linker comprises an oligonucleotide.
- 122. The method according to any of the claims 118 to 121, wherein the oligonucleotide complementing element and the oligonucleotide linker is a continous oligonucleotide.
- 123. The method according to any of the claims 121 to 122, wherein two linkers of distinct building blocks each comprises one part of a molecule pair, said pair being capable of reversible interaction.

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124. The method according to claim 123, wherein the molecule pair comprises a double stranded sequence of nucleic acids forming a dimerisation domain on the linkers.

- 5 125. The method according to claim 123 or 124, wherein the dimerisation domain part of a linker is proximal to the functional entity.
- 126. The method according to claim 124 or 125, wherein the sequence of nucleic acids monomers forming the one part of the molecule pair is separated with 2, 1, or 0 nucleic acid monomers from the functional entity.

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- The method according to claim 126, wherein the functional entity is directly attached to the dimerisation domain.
- 128. The method according to any of the claims 123 to 127, wherein the sequence of nucleic acid monomers comprises 3 to 20 mononucleic acids.

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129. The method according to claim 128, wherein the sequence comprises3 to 15 nucleic acid monomers.

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- The method according to claim 129, wherein the sequence comprises
 4 to 12 nucleic acid monomers.
- 131. The method according to any of the claims 114 to 130, wherein the functional entity is attached to a nucleobase of the oligonucleotide linker.
- 132. The method according to claim 131, wherein the functional entity is attached to a terminal nucleotide of the linker.
- 30 133. The method of claim 1, wherein the complementing elements are selected from nucleotides, and the complementing elements are linked by using a chemical agent, pH change, light, a catalyst, radiation, such as electromagnetic radiation, or by spontaneous coupling when being brought into reactive contact with each other.

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- 134. The method according to claim 133, wherein the chemical agent comprises a phosphoimidazolid group.

The method according to claim 133 and 134, wherein the

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- 5 phospholmidazolid group is attached to the 5' end of the building block and the building block is linked to a nascent complementing template at the 3' hydroxyl group thereof.
- Method of daim 1, wherein at least a subset of said plurality of building
 blocks preferably comprises one complementing element and/or one functional entity and/or one linker.
- 137 The method of claim 1, wherein a subset of said plurality of building blocks comprises a selectively cleavable linker separating the functional entity
 - from the complementing element, wherein said selectively cleavable linker is not deaved under conditions resulting in cleavage of cleavable linkers separating the functional entity from the complementing element of building blocks not belonging to the subset of building blocks comprising a selectively cleavable linker.

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- 138. The method according to claim 102, wherein the functional entity is released by cleaving the cleavable linkers such that no trace of the linker appears on the functional entity.
- cleaved, and wherein the at least one selectively cleavable linkers are cleaved, and wherein the at least one selectively cleavable linker is not cleaved, and wherein the templated molecule is linked to the template and/or to the complementing element by means of said at least one selectively cleavable

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140. The method of claims 138 to 139, wherein the linkers are cleaved by acid, base, a chemical agent, light, electromagnetic radiation, an enzyme, or a catalyst.

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The method according to claim 140, wherein the linker cleaved by light comprises a nitrophenyl moiety. 14

in the range of from $0.8~\mathrm{A}$ to about $5~\mathrm{A}$, for example in the range of from $0.8~\mathrm{A}$ to 25 A, for example in the range of from 0.8 A to about 20 A, such as in the range 0.8 A to about 7 A, for example in the range of from 0.8 A to about 6 A, such as cleavable linker is in the range of from about 0.8 Å to about 70 Å, such as in the about 50 Å, such as in the range of from 0.8 Å to about 40 Å, for example in the of from 0.8 A to about 18 A, for example in the range of from 0.8 A to about 16 Å, such as in the range of from 0.8 Å to about 14 Å, for example in the range of about 4 A, such as in the range of from 0.8 A to about 3.5 A, for example in the range of from 0.8 Å to about 3.0 Å, such as in the range of from 0.8 Å to about range of from 0.8 Å to about 30 Å, such as in the range of from 0.8 Å to about The method of claim 1, wherein the length of the linker or selectively from 0.8 Å to about 12 Å, such as in the range of from 0.8 Å to about 10 Å, for example in the range of from 0.8 A to about 8 A, such as in the range of from range of from 0.8 Å to about 1.5 Å, for example in the range of from 0.8 Å to range of from 0.8 A to about 60 A, for example in the range of from 0.8 A to 2.5 A, for example in the range of from 0.8 A to about 2.0 A, such as in the 42

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30 A, such as in the range of from 1 A to about 25 A, for example in the range of example in the range of from 1 A to about 16 A, such as in the range of from 1 A from 1 A to about 6 A, such as in the range of from 1 A to about 5 A, for example to about 14 Å, for example in the range of from 1 Å to about 12 Å, such as in the range of from 1 A to about 10 A, for example in the range of from 1 A to about 8 in the range of from 1 Å to about 4 Å, such as in the range of from 1.0 Å to abou range of from 1 Å to about 40 Å, for example in the range of from 1 Å to about The method of claim 1, wherein the length of the linker or selectively cleavable linker is in the range of from about 1 A to about 60 A, such as in the A, such as in the range of from 1 A to about 7 A, for example in the range of range of from 1.0 Å to about 2.5 Å, for example in the range of from 1.0 Å to 3.5 A, for example in the range of from 1.0 A to about 3.0 A, such as in the from 1 Å to about 20 Å, such as in the range of from 1 Å to about 18 Å, for

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about 2.0 A, such as in the range of from 1.0 A to about 1.5 A, for example in the range of from 1.0 Å to about 1.2 Å.

2 A to about 18 A, for example in the range of from 2 A to about 16 A, such as in range of from 2 Å to about 8 Å, such as in the range of from 2 Å to about 7 Å, for range of from 2 Å to about 30 Å, such as in the range of from 2 Å to about 25 Å, for example in the range of from 2 Å to about 20 Å, such as in the range of from example in the range of from 2 A to about 6 A, such as in the range of from 2 A cleavable linker is in the range of from about 2 A to about 40 A, such as in the about 12 A, such as in the range of from 2 A to about 10 A, for example in the The method of claim 1, wherein the length of the linker or selectively the range of from 2 A to about 14 A, for example in the range of from 2 A to 4. 2

about 3.0 Å, such as in the range of from 2.0 Å to about 2.5 Å, for example in the to about 5 A, for example in the range of from 2 A to about 4 A, such as in the range of from 2.0 A to about 3.5 A, for example in the range of from 2.0 A to range of from 2.0 A to about 2.2 A.

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4 A to about 18 A, for example in the range of from 4 A to about 16 A, such as in range of from 4 Å to about 8 Å, such as in the range of from 4 Å to about 7 Å, for range of from 4 $^{\rm A}$ to about 30 Å, such as in the range of from 4 Å to about 25 Å, for example in the range of from 4 Å to about 20 Å, such as in the range of from example in the range of from 4 Å to about 6 Å, such as in the range of from 4 Å cleavable linker is in the range of from about 4 Å to about 40 Å, such as in the about 12 A, such as in the range of from 4 A to about 10 A, for example in the The method of claim 1, wherein the length of the linker or selectively the range of from 4 A to about 14 A, for example in the range of from 4 A to to about 5 A. 145

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about 1.0 A.

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6 A to about 18 A, for example in the range of from 6 A to about 16 A, such as in for example in the range of from 6 A to about 20 A, such as in the range of from range of from 6 A to about 30 A, such as in the range of from 6 A to about 25 A, cleavable linker is in the range of from about 6 Å to about 40 Å, such as in the The method of claim 1, wherein the length of the linker or selectively the range of from 6 A to about 14 A, for example in the range of from 6 A to

about 12 A, such as in the range of from 6 A to about 10 A, for example in the range of from 6 A to about 8 A, such as in the range of from 6 A to about 7 A.

8 A to about 18 A, for example in the range of from 8 A to about 16 A, such as in for example in the range of from 8 A to about 20 A, such as in the range of from range of from 8 Å to about 30 Å, such as in the range of from 8 Å to about 25 Å, The method of claim 1, wherein the length of the linker or selectively cleavable linker is in the range of from about 8 Å to about 40 Å, such as in the the range of from 8 A to about 14 A, for example in the range of from 8 A to about 12 Å, such as in the range of from 8 Å to about 10 Å.

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means of a reaction involving reaective groups, a functional group of at least one The method according to claim 1, whereinthe obtained templated molecule comprises a sequence of covalently linked functional groups, by functional entity to a functional group of a neighbouring functional entity.

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- The method according to any of the claims 1 to 148, wherein the templated molecule is a linear sequence of functional groups. 149.
- The method according to any of the claims 1 to 148, wherein the templated molecule is a branched sequence of functional groups. 50. 20
- The method according to any of the claims 1 to 148, wherein the templated molecule is a cyclic sequence of functional groups. 51.
- The method of claim 1, wherein the templated molecule is an oligomer or a polymer comprising at least one repetitive sequence of functional groups. 152

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- Method of claim 152, wherein the sequence of at least three functional groups is repeated at least twice in the templated molecule 53 ဓ္က
- Method of claim 152, wherein any sequence of at least three functional groups in the templated molecule occurs only once. 54.

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- The method of claim 1, wherein the templated molecule comprises or essentially consists of amino acids selected from the group consisting of $\alpha\text{-}$ amino acids, β-amino acids, γ-amino acids, ω-amino acids.
- The method of claim 1, wherein the templated molecule comprises or essentially consists of natural amino acid residues. 156. S
- The method of claim 1, wherein the templated molecule comprises or essentially consists of a-amino acids. 157.

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- The method of claim 1, wherein the templated molecule comprises or essentially consists of monosubstituted a-amino acids. 158.
- The method of claim 1, wherein the templated molecule comprises or essentially consists of disubstituted α -amino acids. 160

The method of claim 1, wherein the templated molecule comprises or

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The method of claim 1, wherein the templated molecule comprises or essentially consists of monosubstituted β-amino acids. 161. ឧ

essentially consists of disubstituted β-amino acids.

- The method of claim 1, wherein the templated molecule comprises or essentially consists of trisubstituted p-amino acids. 162.
- The method of claim 1, wherein the templated molecule comprises or essentially consists of tetrasubstituted β-amino acids. 163.

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The method of any of claims 115 to 118, wherein the backbone structure of said B-amino acids comprises or essentially consists of a cyclohexane-backbone and/or a cyclopentane-backbone 蝥

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The method of claim 1, wherein the templated molecule comprises or essentially consists of y-amino acids. 165.

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166. The method of claim 1, wherein the templated molecule comprises or essentially consists of \(\partial\)-amino acids.

 The method of claim 1, wherein the templated molecule comprises or essentially consists of vinylogous amino acids.

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168. The method of claim 1, wherein the templated molecule comprises or essentially consists of N-substituted glycines.

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essentially consists of molecules or molecular entities selected from the group of polypeptidylphosphonates, polyurethanes, azatides, oligo N-substituted glycines, The method of claim 1, wherein the templated molecule comprises or α -peptides, β -peptides, γ -peptides, ω -peptides, mono-, di- and tri-substituted α residues are in the L-form or in the D-form, vinylogous polypeptides, glycopolypeptides, B-peptides, y-peptides, w-peptides, peptides wherein the amino acid polyethers, ethoxyformacetal oligomers, poly-thioethers, polyethylene glycols polyvinyl, lipids, phospholipids, glycolipids, polycyclic compounds comprising (PEG), polyethylenes, polydisulfides, polyarylene sulfides, polynucleotides, peptides, polyamides, vinylogous sulfonamide peptide, polysulfonamide, polyethyleneimines, polyimides, polyacetals, polyacetates, polystyrenes, a.g. aliphatic or aromatic cycles, including polyheterocyclic compounds, PNAs, LNAs, morpholinos, oligo pyrrolinone, polyoximes, polyimines, proteoglycans, and polysiloxanes, inlcuding any combination thereof. conjugated peptides comprising e.g. prosthetic groups, polyesters, polysaccharides, polycarbamates, polycarbonates, polyureas, 169

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170. The method of claim 1, wherein neighbouring residues of the templated molecule is linked by a chemical bond selected from the group of chemical bonds consisting of peptide bonds, sulfonamide bonds, ester bonds, saccharide bonds, carbamate bonds, carbonate bonds, urea bonds, phosphonate bonds, urea bonds, carbonate bonds, peptoid bonds, ether bonds, ethoxy bonds, thioether bonds, single carbon bonds, disulfide bonds, sulfide bonds, double carbon bonds, triple carbon bonds, disulfide bonds, sulfide bonds, phosphodiester bonds, oxime bonds, imine bonds, imide bonds, including any combination thereof.

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171. The method according to claim 170, wherein the chemical bond linking functional entities of two or more building blocks is formed by a reaction of a nucleophile group of a first functional entity with an ester or thioester of another functional entity.

172. The method according to claim 171, wherein the linker of the building block bearing the thioester group is cleaved simultaneously with the formation of the bonding resulting in a transfer of functional group to the nucleophilic building block.

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- 173. The method according to claim 171 and 172, wherein the nucleophile group is selected from -NH₂, H₂NHN, HOHN-, H₂N-C(O)-NH-.
- 15 The method of claim 1, wherein the backbone structure of said templated molecule comprises or essentially consists of a molecular group selected from -NHN(R)CO-; -NHB(R)CO-; -NHC(RR)CO-; -NHC(RR)CO-; -COCH₂-; -COS-; -CONR-; -CHO-; -CH₂-CH₂-CH₂-; -COS-; -CONR-; -CH₂-CH₂-CH₂-CH₂-; -CH₂-
- 20 CH(CH₃)S-;-CH=CH-;-NHCO-;-NHCONH-;-CONHO-;-C(=CH₂)CH₂-;-PO₂NH-;-PO₂CH₂-;-PO₂CH₂N²-;-SO₂NH-; and lactams.
- 175. The method of claim 1, wherein the functional entity is selected from the group of precursors selected from α-amino acid precursors, β-amino acid precursors, γ-amino acid precursors, and α-amino acid precursors.

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- 176. The method of claim 175, wherein the β-amino acid precursor is a N-carboxyanhydride ring structure derivative.
- 30 177. The method of claim 171, wherein the β-amino acid precursor is an thiocarboxyanhydrid (NTA) ring structure derivative.
- 178. The method according to claim 1, wherein a functional group af at least one functional entity is linked to another functional entity by means of a bridging molecule.

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179. The method according to claim 178, wherein the functional entities are neighbours.

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180. The method of claim 1, wherein the templated molecule comprises or essentially consists of at least 2 different functional groups, such as at least 3 different functional groups, for example at least 4 different functional groups, such as at least 5 different functional groups, for example at least 6 different functional groups, such as at least 7 different functional groups, for example at least 8 different functional groups, such as at least 9 different functional groups, for example at least 10 different functional groups, for example at least 10 different functional groups.

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181. The method of claim 180, wherein the functional groups are identical.

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- 182. The method of claim 1, wherein each building block comprises at least one reactive group type I.
- 183. The method according to claim 1, wherein each building block comprises at least one reactive group type II.

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184. The method of claim 182, wherein each building block comprises two

reactive groups type I.

- 25 185. The method of claim 183, wherein each building block comprises two reactive groups type II.
- 186. The method according to claim 185, wherein at least one building block comprises three or more, such as four or more reactive groups type II.
- The method according to daims 185 or 186, wherein the reactive groups type II are similar.

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The method according to claims 185 or 186, wherein the reactive groups type II are reaction partners.

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189. The method according to claims 187 or 188, wherein the reactive groups type II are capable of forming a linkage between the functional groups through a bridging molecule.

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190. The method according to claim 189, wherein the bridging molecule is a di-functional compound with reactive groups which are reaction partners to the reactive groups type II of the functional entities.

10 191. The method according to claim 185, wherein the reactive groups type Il are non-similar.

192. The method according to claim 191, wherein the non-similar reactive groups type if are reaction partners.

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193. The method according to claim 191 or 192, wherein the reactive groups type II are capable of forming a linkage between the functional groups through a bridging molecule. 194. The method according to claim 191, wherein the bridging molecule is a di-functional compound with reactive groups which are reaction partners to the reactive groups type II of the functional entities.

195. The method of any of claims 182 to 194, wherein the at least one reactive group type II of the functional entity is selected from the group consisting of N-carboxyanhydride (NCA), N-thiocarboxyanhydride (NTA), coumarin, amine, carboxylic acid, ketone, aldehyde, hydroxyl, thiol, ester, thioester, alkenyl, any conjugated system of double bonds, hydrazine, N-hydroxysuccinimide ester, and epoxide.

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 The method of claims 178 to 190, wherein the reactive group type II is an electrophile.

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197. The method according to claim 191, wherein the reactive group type II
 35 is an electrophile selected from the group consisting of -X-C(O)-R, -X-C(O)-

CHR-C(O)-R, -X-C(O)-CR=CH-R, and -X-C(O)-CHR-NHR, wherein X is a S or O, and R independently is a functional group. The method of claims 183 to 195, wherein the reactive group type II is a nucleophile. 198.

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- The method according to claim 198, wherein the reactive group type II is a nucleophile selected from the group consisting of -NH2, H2NHN-, HOHN-, and H₂N-C(0,S)-NH-. 199.
- The method of claims 183 to 195, wherein the reactive group type II is a radical. 500 500

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- The method according to any of the preceding claims, wherein the functional entity comprises a reactive group type II connected to the linker. 2 5
- The method according to claim 201, wherein said functional entity firther comprises at least one further reactive group type II. 202
- comprising the at least one further reactive group type II is capable of forming a between said functional entity and the linker, by a reaction involving the reactive link to a second functional entity having a reactive group type II interspaced The method according to claim 202, wherein the functional entity group type II on said other functional entity. 203

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the functional group of the second functional entity to the first functional entity. second functional entity and the linker is cleaved resulting in a translocation of The method according to claim 203, wherein the link between the 8

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linkage between the first and the second functional entity is formed by a direct The method according to any of the claims 201 to 204, wherein the reaction of the respective reactive groups type II. 8 ၉

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- The method according to any of the claims, 201 to 204, wherein the linkage between the first and the second functional entity is formed through a bridging molecule.
- The method according to any of the preceding claims, wherein multiple functional groups are linked by a reaction cascade involving a plurality of building blocks. 207.
- multiple functional groups are linked by a reaction cascade involving a plurality of building blocks, each of said building blocks comprising a functional entity which comprises a first reactive group type II connected to the linker, said The method according to any of the claims 201 to 207, wherein functional entity comprising at least one further reactive group type II. 208

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- functional entity is restricted as to rotation to obtain a preferred orientation during The method according to any of the preceding claims, wherein the formation of linkages to other functional entitles. 209 5
- that links the functional entity to the complementing element is fixed with regard The method according to claim 209, wherein one or more bondings to rotation. 210.

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preferred orientation is obtained by coupling the functional entity to the base as The method according to any of the claims 209 and 210, wherein the well as the (deoxy)ribose moiety of a nucleotide building block. 211.

- The method according to any of the claims 209 to 211, wherein the functional entity is coupled to two bases of a dinucleotide. 212.
- formation of the templated molecule involves the reactive groups type II capable The method according to any of the preceding claims, wherein the of forming a polymer through anionic, cationic, or radical polymerisation. 213 ဓ
- The method according to any of the preceding claims, comprising a scaffold functional entity comprising one of more reactive groups type II, said 214. 33

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scaffold functional entity being the basic chemical structure for forming a templated molecule by addition of functional groups emanating from the plurality of building blocks.

- 5 215. The method according to claim 2014, wherein the scaffold functional entity comprises two or more reactive groups type II.
- 216. The method according to claim 214 or 215, wherein the scaffold functional entity is linked covalently to the template.

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- 217. The method according to any of the claims 214 to 215, wherein the scaffold functional entity is linked to a complementing element capable of recognising a coding element of the template.
- The method according to any of the claims 214 to 217, wherein building blocks attached to the template comprises functional entities carrying reactive groups type II capable of forming a link to the scaffold functional entity
- 219. The method according to any of the claims 214 to 218, wherein a scrambling of linking of functional groups to the scaffold functional entity is obtained by having a different number of reactive groups type II on the scaffold functional entity compared to the number of reactive groups type II on the functional entities of the plurality of building blocks.

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- 25 220. The method according to any of the daims 214 to 219, wherein the reaction of the reactive groups type II on the scaffold and the corresponding reactive groups type II on the building blocks result in a simultaneous linkage of the functional groups to the scaffold functional entity and deavage of the linker connecting the functional entity with the complementing element.
- 221. The method according to any of the preceding claims, wherein a plurality of templates having different coding elements and/or different order of coding elements is used.

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222. The method according to claim 221, wherein two or more different templates are used.

The method according to claim 221 or 222, wherein four or more

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- 5 different templates, such as more than 103, 105, 107, 109, 1011, 1013, 1015, 1017 different templates are used.
- 224. The method according to claim 221, wherein the plurality of different templates results in a library of different templated molecules, each of said templated molecules being connected to the specific template or complementing

template that templated the molecule.

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225. The method of any of the preceding claims comprising the further step of releasing the template or complementing template from the templated molecule, and obtaining a templated molecule that is not linked to the complementing template or template that templated the synthesis of the templated molecule.

- 26. A templated molecule obtainable in accordance with any of the claims
 1 to 225, a plurality of covalently linked functional groups each comprising a
 residue, wherein the covalently linked residues are capable of generating a
 polymer comprising, exclusively or in combination with additional portions, at
 least one portion selected from the group of polymer portions consisting of αpeptides, β-peptides, γ-peptides, α-peptides, mono-, di- and tri-substituted αpeptides, β-peptides, γ-peptides, α-peptides, peptides wherein the amino acid
 residues are in the L-form or in the D-form, vinylogous polypeptides, glycopolypeptides, polyamides, vinylogous sulfonamide peptides, polysulfonamides,
 conjugated peptides comprising e.g. prosthetic groups, polysesters,
 polysaccharides, polycarbamates, polycarbonates, polyureas,
- polypeptidylphosphonates, polyurethanes, azatides, oligo N-substituted glycines, polyethers, ethoxyformacetal oligomers, poly-thioethers, polyethylene glycols (PEG), polyethylenes, polydisulfides, polyarylene sulfides, polynucleotides, PNAs, LNAs, morpholinos, oligo pyrrolinones, polyoximes, polyimines, polyethyleneimines, polyimides, polyacetals, polyacetates, polystyrenes,
 - 35 polyvinyl, lipids, phospholipids, glycolipids, polycyclic compounds comprising

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rom 2 to 20, for example from 2 to 15, such as from 2 to 10, such as from 2 to 8, o 8, such as from 4 to 6, for example 4, for example from 5 to 100, such as from 12 to 40, for example from 12 to 30, such as from 12 to 20, for example from 12 rom 3 to 15, such as from 3 to 10, such as from 3 to 8, for example from 3 to 6, such as from 3 to 4, for example 3, such as from 4 to 100, for example from 4 to as from 5 to 8, for example from 5 to 6, for example 5, such as from 6 to 100, for 10, such as from 6 to 8, such as 6, for example from 7 to 100, such as from 7 to example 9, for example from 10 to 100, such as from 10 to 80, for example from to 15, such as from 14 to 100, such as from 14 to 80, for example from 14 to 60, from 12 to 100, such as from 12 to 80, for example from 12 to 60, such as from as from 4 to 20, such as from 4 to 15, for example from 4 to 10, such as from 4 30, such as from 4 to 60, such as from 4 to 40, for example from 4 to 30, such from 6 to 30, such as from 6 to 20, such as from 6 to 15, for example from 6 to rom 7 to 8, for example 7, for example from 8 to 100, such as from 8 to 80, for example from 8 to 60, such as from 8 to 40, for example from 8 to 30, such as 10 to 60, such as from 10 to 40, for example from 10 to 30, such as from 10 to example from 16 to 60, such as from 16 to 40, for example from 16 to 30, such example from 2 to 60, such as from 2 to 40, for example from 2 to 30, such as example from 3 to 30, such as from 3 to 20, such as from 3 to 15, for example 5 to 80, for example from 5 to 60, such as from 5 to 40, for example from 5 to 30, such as from 5 to 20, for example from 5 to 15, such as from 5 to 10, such example from 6 to 80, such as from 6 to 60, such as from 6 to 40, for example 20, for example from 10 to 15, such as from 10 to 12, such as 10, for example such as from 7 to 20, for example from 7 to 15, such as from 7 to 10, such as or example from 2 to 6, such as from 2 to 4, for example 2, such as from 3 to 100, for example from 3 to 80, such as from 3 to 60, such as from 3 to 40, for 80, for example from 7 to 60, such as from 7 to 40, for example from 7 to 30, preferably from 2 to 200, for example from 2 to 100, such as from 2 to 80, for such as from 14 to 40, for example from 14 to 30, such as from 14 to 20, for rom 8 to 20, for example from 8 to 15, such as from 8 to 10, such as 8, for example from 14 to 16, such as from 16 to 100, such as from 16 to 80, for proteoglycans, and polysiloxanes, and wherein the plurality of residues is a.g. aliphatic or aromatic cycles, including polyheterocyclic compounds, S 2 5 22 ဓ 8

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60, such as from 35 to 40, for example from 40 to 100, such as from 40 to 80, for as from 45 to 100, for example from 45 to 80, such as from 45 to 60, for example to 20, for example from 20 to 100, such as from 20 to 80, for example from 20 to 60 to 80, such as from 60 to 70, for example from 70 to 100, such as from 70 to 60, such as from 20 to 40, for example from 20 to 30, such as from 20 to 25, for example from 22 to 100, such as from 22 to 80, for example from 22 to 60, such from 45 to 50, such as from 50 to 100, for example from 50 to 80, such as from as from 22 to 40, for example from 22 to 30, such as from 22 to 25, for example from 25 to 100, such as from 25 to 80, for example from 25 to 60, such as from example from 40 to 60, such as from 40 to 50, for example from 40 to 45, such 30 to 80, such as from 30 to 60, for example from 30 to 40, such as from 30 to 35, for example from 35 to 100, such as from 35 to 80, for example from 35 to 50 to 60, for example from 50 to 55, such as from 60 to 100, for example from 90, for example from 70 to 80, such as from 80 to 100, for example from 80 to 25 to 40, for example from 25 to 30, such as from 30 to 100, for example from 90, such as from 90 to 100. \$

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The templated molecule according to claim 226, wherein the templated molecule is a polymer. 227.

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The templated molecule according to claim 227, wherein the polymer is linear. 228.

The temnplated molecule according to claim 227, wherein the polymer 229.

is branched. 22

exclusively or in combination with additional portions selected from the group, at least one portion selected from the group of polymer portions consisting of $\alpha\text{-}$ peptides, β-peptides, γ-peptides, ω-peptides, peptides wherein the amino acid The templated molecule according to claim 226 or 229, wherein the peptides, B-peptides, γ -peptides, ω -peptides, mono-, di- and tri-substituted α covalently linked residues are capable of generating a polymer comprising, residues are in the L-form or in the D-form, and vinylogous polypeptides. 330

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from 18 to 60, such as from 18 to 40, for example from 18 to 30, such as from 18

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as from 16 to 20, such as from 18 to 100, such as from 18 to 80, for example

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- The templated motecule according to claims 226 to 230, wherein the covalently linked residues are capable of generating a polysaccharaide.
- 232. A templated molecule according to any of the claims 226 to 231, comprising a sequence of functional groups, wherein neighbouring functional groups are linked by a molecular molety that is not natively associated with said functional groups.

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233. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule does not comprise or consist of an α-peptide or a nucleotide.

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234. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule does not comprise or consist of a monosubstituted α-peptide or a nucleotide.

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235. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule does not comprise or consist of a peptide or a nucleotide.

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any of the claims 221 to 224, wherein said composition comprises a plurality of more than or about 10³ different templated molecules, such as more than or about 10⁴ different templated molecules, for example more than or about 10⁴ different templated molecules, for example more than or about 10⁵ different templated molecules, such as more than or about 10⁵ different templated molecules, such as more than or about 10⁷ different templated molecules, for example more than or about 10¹⁰ different templated molecules, such as more than or about 10¹⁰ different templated molecules, such as more than or about 10¹⁰ different templated molecules, such as more than or about 10¹⁰ different templated molecules, such as more than or about 10¹⁰ different templated molecules, for example more than or about 10¹⁰ different templated molecules, for example more than or about 10¹⁰ different templated molecules, such as more than or about 10¹⁰ different templated molecules, so as more than or about 10¹⁰ different templated molecules, for example

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more than or about 10^{17} different templated molecules, such as more than or about 10^{18} different templated molecules.

- 237. The composition according to claim 236, wherein the plurality of templated molecules is selected from the group of templated molecules defined in any of the claims 226 to 235.
- 238. The composition according to any of claims 236 and 237, wherein said composition further comprises the template capable of templating each

templated molecule, or a subset thereof.

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- 239. A composition according to claim 238, wherein the templated molecule is linked to the template capable of templating the templated molecule.
- 240. The composition according to claim 239, wherein the template motecule is covalently linked to the template.
- 241. The composition according to claim 239, wherein the templated molecule is linked non-valently to the template.

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- 242. The composition according to claim 241, wherein the non-covalent linkage involves one or more bondings selected from the group consisting of hydrogen bondings, van der Waals bondings, and ionic bondings.
- 25 243. A complex comprising a templated molecule and the template that templated the synthesis of the templated molecule.
- 244. The composition according to any of claims 236 to 242, wherein the template is not a natural nucleotide, when the templated molecule is an α-

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245. The composition according to any of claims 236 to 242, wherein the templated molecule is not a naturally occurring α-peptide.

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246. The composition according to any of claims 236 to 242, wherein the templated molecule is not an α-peptide.

- The composition according to claim 236 to 242, wherein the template
 does not consist exclusively of natural nucleotides, when the templated molecule
 is a peptide comprising exclusively monosubstituted α-amino acids.
- 248. The composition according to claim 236 to 242, wherein the template is not a natural nucleotide, when the templated molecule is a natural α-peptide.

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- 249. The composition according to claim 236 to 242, wherein the template is not a nucleotide, when the templated molecule is a natural α-peptide.
- The composition according to claim 236 to 242, wherein the template
 is not a nucleotide, when the templated molecule is a monosubstituted α-peptide.

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 The composition according to claim 236 to 242, wherein the template is not a nucleotide, when the templated molecule is an α-peptide.

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- 252. The composition according to claim 236 to 242, wherein the template is not a natural nucleotide, when the templated molecule is a peptide.
- The composition according to claim 236 to 242, wherein the template
 is not a nucleotide, when the templated molecule is a peptide.
- 254. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the templated molecule does not comprise or consist of an α-peptide
- 255. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the

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templated molecule, wherein the templated molecule does not comprise a monosubstituted $\alpha\text{-peptide}.$

- 256. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the templated molecule does not comprise or consist of an α-peptide or a nucleotide.
- 257. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template is not a natural nucleotide, when the templated molecule is an α-peptide.

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258. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template does not consist exclusively of natural nucleotides, when the templated molecule is a peptide comprising exclusively monosubstituted α-amino acids.

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- 259. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template is not a natural nucleotide, when the templated molecule is a natural α-peptide.
- 260. A templated molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the templated molecule, wherein the template is not a nucleotide, when the templated molecule is a natural o-peptide.

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functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the A templated molecule comprising a sequence of covalently linked, templated molecule, wherein the template is not a nucleotide, when the templated molecule is a monosubstituted α-peptide. 261.

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functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the A templated molecule comprising a sequence of covalently linked, templated molecule, wherein the template is not a nucleotide, when the templated molecule is an α-peptide.

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functional groups, wherein the templated molecule is linked by means of a linker templated molecule, wherein the template is not a natural nucleotide, when the to the complementing template or template that templated the synthesis of the A templated molecule comprising a sequence of covalently linked, templated molecule is a peptide. . 283

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functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that templated the synthesis of the A templated molecule comprising a sequence of covalently linked, templated molecule, wherein the template is not a nucleotide, when the templated molecule is a peptide. 264

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The templated molecule according to any of claims 226 to 235, and 254 to 264, wherein the templated molecule is an oligomer or a polymer comprising at least one repetitive sequence of functional groups. 265 22

254 to 264, wherein the sequence of at least three functional groups is repeated The templated molecule according to any of claims 226 to 235, and at least twice in the templated molecule. 266.

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The templated molecule according to any of claims 226 to 235, and 254 to 264, wherein any sequence of at least three functional groups in the templated molecule occurs only once. 267.

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254 to 264, wherein the templated molecule comprises or essentially consists of amino acids selected from the group consisting of lpha-amino acids, eta-amino acids, The templated molecule according to any of claims 226 to 235, and . 288

γ-amino acids, ω-amino acids. သ

254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and natural amino acid residues. 269

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254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and a-amino acids.

254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and monosubstituted α -amino acids. 271. ₹

254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and disubstituted α-amino acids 272. ន

254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and 273.

monosubstituted β-amino acids. 22

254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and disubstituted \(\theta\)-amino acids. 274.

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254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and trisubstituted \(\beta\)-amino acids. 275.

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254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and tetrasubstituted β-amino acids. 276.

backbone structure of said β -amino acids comprises or essentially consists of a The templated molecule of any of claims 273 to 276, wherein the cyclohexane-backbone and/or a cyclopentane-backbone. 277.

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- 254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and y-amino acids. 278. 2
- 254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and w-amino acids. 279.

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254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and vinylogous amino acids. **5**80.

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- 254 to 264, wherein the templated molecule comprises or essentially consists of The templated molecule according to any of claims 226 to 235, and N-substituted glycines. 281.
- 254 to 264, wherein the templated motecule comprises or essentially consists of peptides, y-peptides, w-peptides, peptides wherein the amino acid residues are The templated molecule according to any of claims 226 to 235, and peptides, y-peptides, e-peptides, mono-, di- and tri-substituted a-peptides, βin the L-form or in the D-form, vinylogous polypeptides, glycopoly-peptides, polyamides, vinylogous sulfonamide peptide, polysulfonamide, conjugated molecules or molecular entities selected from the group of $\alpha\text{-peptides},\ \beta\text{-}$ ethoxyformacetal oligomers, poly-thioethers, polyethylene glycols (PEG), peptides comprising e.g. prosthetic groups, polyesters, polysaccharides, polycarbamates, polycarbonates, polyureas, polypeptidylphosphonates, polyurethanes, azatides, oligo N-substituted glycines, polyethers, 52 ဓ္က

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polyvinyl, lipids, phospholipids, glycolipids, polycyclic compounds comprising polyethylenes, polydisulfides, polyarylene sulfides, polynucleotides, PNAs, polyethyleneimines, polyimides, polyacetals, polyacetates, polystyrenes, e.g. aliphatic or aromatic cycles, including polyheterocyclic compounds, proteoglycans, and polysiloxanes, inlcuding any combination thereof. LNAs, morpholinos, oligo pyrrolinone, polyoximes, polyimines,

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single carbon bonds, double carbon bonds, triple carbon bonds, disulfide bonds, sulfide bonds, phosphodiester bonds, oxime bonds, imine bonds, imide bonds, peptide bonds, sulfonamide bonds, ester bonds, saccharide bonds, carbamate 254 to 264, wherein neighbouning residues of the templated molecule is linked The templated molecule according to any of claims 226 to 235, and by a chemical bond selected from the group of chemical bonds consisting of bonds, carbonate bonds, urea bonds, phosphonate bonds, urethane bonds, azatide bonds, peptoid bonds, ether bonds, ethoxy bonds, thioether bonds, amine, amide, including any combination thereof.

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The templated molecule according to claim 283, wherein the templated molecule is not an a-peptide 284.

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The templated molecule according to any of claims 226 to 235, and -NHC₆ H₄ CO-; -NHCH₂ CHRCO-; -NHCHRCH₂ CO-; -COCH₂-; -COS-; -CONR-; -COO-; -CSNH-; -CH2 NH-; -CH2CH2-; -CH2 S-; -CH2 SO-; -CH₂SO₂-; -CH(CH₃)S-; -CH=CH-; -NHCO-; -NHCONH-; -CONHO-; 254 to 264, wherein the backbone structure of said templated molecule comprises or essentially consists of a molecular group selected from -NHN(R)CO-; -NHB(R)CO-; -NHC(RR')CO-; -NHC(=CHR)CO-; 22

-C(=CH₂)CH₂-; -PO₂·NH-; -PO₂·CH₂-; -PO₂·CH₂N'-; -SO₂NH'-; and lactams,

including any combination thereof.

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selected from α -amino acid precursors, eta-amino acid precursors, γ -amino acid The temptated motecule according to any of claims 226 to 235, and 254 to 264, wherein the precursor is selected from the group of precursors precursors, and w-amino acid precursors. 286.

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287. The templated molecule according to any of claims 219 to 226, and 240 to 250, wherein the templated molecule comprises or essentially consists of at least 2 different functional groups, such as at least 3 different functional groups, for example at least 4 different functional groups, such as at least 5 different functional groups, for example at least 6 different functional groups, such as at least 7 different functional groups, for example at least 8 different functional groups, such as at least 9 different functional groups, for example at least 10 different functional groups.

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 The templated molecule according to claim 287, wherein the functional groups are identical.

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289. The templated molecule according to any of claims 226 to 235, and 254 to 264, wherein the templated molecule is obtainable by a method according to any of claims 1 to 218.

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290. A molecule comprising a sequence of covalently linked building blocks, wherein the sequence of covalently linked building blocks comprises a sequence of complementing elements forming a complementing template capable of complementing the template that templated the synthesis of the templated molecule, and wherein the templated molecule is linked to the complementing template or template that templated its synthesis.

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291. A templated molecule according to any of the previous claims, wherein the templated molecule comprises a sequence of functional entities comprising at least one functional group, and optionally at least one reactive group type II, and wherein each functional entity is linked to a complementing element or a template that templated the synthesis of the templated molecule.

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292. The composition according to claims 226 to 291, wherein the templated is prepared in accordance with claim 1 to 225, where said templated molecule is bound to another molecule selected from the group consisting of DNA, RNA, antibody, peptide, or protein, or derivatives thereof.

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293. A method for screening templated molecules potentially having a predetermined activity, said method comprising the step of providing a target molecule or a target entity, including a surface, and obtaining templated molecules having an affinity for - or an effect on - said target molecule or target

entity.

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294. A method for assaying an activity potentially associated with a templated molecules, said method comprising the step of providing a target molecule or a target entity, including a surface, and obtaining templated molecules having an affinity for - or an effect on - said target molecule or target entity, and determining the activity of the templated molecule.

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295. A method for selecting complexes or templated molecules having a predetermined activity, said method comprising the step of performing a selection procedure and selecting templated molecules based on predetermined selection criteria.

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 A method for screening a composition of molecules having a predetermined activity comprising:

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 establishing a first composition of templated molecules, sald molecules being defined as in any of the claims 226 to 235, and 254 to 290, or produced as defined in any of the claims 1 to 225, ii) exposing the first composition to conditions enriching said first composition with templated molecules having the predetermined activity, and

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 iii) optionally amplifying the templated molecules of the enriched composition obtaining a second composition,

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(v) further optionally repeating step ii) to iii), and

 obtaining a further composition having a higher ratio of templated molecules having the specific predetermined activity.

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templated molecules, wherein said mutagenesis can take place prior to carrying out step iil), simultaneously with carrying out step iil), or after carrying out step The method of claim 296, further comprising a step of mutating the 297.

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The method of claim 297, wherein the mutagenesis is carried out as random or site-directed mutagenesis. 298.

The method of claim 296, wherein step iii) comprises a 101 to 1015-fold amplification. 299. 2

The method of claim 296, wherein the steps ii) and iii) are repeated at least 2, 3, 5 times, such as at least 10 times, such as at least 15 times. g S

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The method of claim 296, further comprising a step of identification of the templated molecule having the predetermined activity. 301.

The method of claim 296, wherein the identification is conducted by analysing the template and/or complementary template physically or by other 305

means associated with the molecule.

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having the predetermined activity, said binding partner being directly or indirectly composition comprises providing a binding partner to said templated molecule The method of claim 296, wherein the conditions enriching the immobilised on a support. 33

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composition involves any one or more of electrophoretic separation, gelfiltration, immunoprecipitation, isoelectric focusing, centrifugation, and immobilization. The method of claim 296, wherein the conditions enriching the 384

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The method of claim 296, wherein the predetermined activity is an enzymatic activity or a catalytic activity. 305.

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molecule, or performing an interaction with the templated molecule having the composition comprises providing cells capable of internalising the templated The method of claim 296, wherein the conditions enriching the predetermined activity. . 8

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A method for amplifying the complementing template or the template having a predetermined activity, said method comprising the step of contacting that templated the synthesis of the templated molecule having, or potentially the template with amplification means, and amplifying the template. 307.

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A method for amplifying the complementing template or the template contacting the template with amplification means, and amplifying the template, that templated the synthesis of the templated molecule having, or potentially and ii) obtaining the templated molecule in an at least two-fold increased having, a predetermined activity, said method comprising the steps of i) 308.

amount.

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A method for altering the sequence of a templated molecule, including generating a templated molecule comprising a novel or altered sequence of functional groups, wherein said method preferably comprises the steps of 309.

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plementing templates or first templates capable of templating a plurality providing a first complementing template or a first template capable of templating the first templated molecule, or a plurality of such first com-=

of first templated molecules,

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mutating or modifying the sequence of the first complementing template or the first template, or the plurality of first complementing templates or first templates, and generating a second template or a second comple-≘

menting template, or a plurality of second templates or second complementing temptates,

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rality of second templated molecules,

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ble of templating the synthesis of a second templated molecule, or a plu-

wherein said second template(s) or complementing template(s) is capa-

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covalently linked, functional groups that is not identical to the sequence of functional groups of the first templated molecule(s), and optionally

wherein said second templated molecule(s) comprises a sequence of

plate(s) a second templated molecule, or a plurality of such second temtemplating by means of said second template(s) or complementing templated molecules. €

A method for altering the sequence of a templated molecule, including generating a templated molecule comprising a novel or altered sequence of 310

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functional groups, wherein said method preferably comprises the steps of

providing a plurality of first complementing templates or first templates capable of templating a plurality of first templated molecules,

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plates or first templates, and generating a second template or a second recombining the sequences of the plurality of first complementing temcomplementing template, or a plurality of second templates or second complementing templates, e

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ble of templating the synthesis of a second templated molecule, or a pluwherein said second template(s) or complementing template(s) is caparality of second templated molecules,

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covalently linked, functional groups that is not identical to the sequence wherein said second templated molecule(s) comprises a sequence of of functional groups of the first templated molecule(s), and optionally

plate(s) a second templated molecule, or a plurality of such second temtemplating by means of said second template(s) or complementing templated molecules. Ê

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The method of any of claims 309 and 310, comprising the further step of amplifying the complementing template or the template that templated the 31.

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synthesis of the templated molecule, said amplification step taking place prior to, simultaneously with, or after the step of mutagenesis or recombination.

- The method of claim 309 to 310, wherein the mutagenesis is 312
- mutagenesis, unique site-elimination (USE), error-prone PCR, error-prone DNA conducted as site-directed mutagenesis, cassette mutagenesis, chemical S
- The method of claim 309 to 310, wherein the mutagenesis is 313
- conducted as DNA shuffling or any form of recombination including homologous recombination either in vivo or in vitro. 2
- A building block comprising 314.

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- element having a recognition group, said complementing element being a complementing element capable of specifically recognising a coding selected from nucleotides, amino acids, antibodies, antigens, proteins, peptides, and molecules with nucleotide recognizing ability, =
- peptides, β-peptides, γ-peptides, ω-peptides, peptides wherein the amino acid residues are in the L-form or in the D-form, vinylogous polypeptides, at least one functional entity selected from a precursor of α -peptides, β peptides, γ -peptides, ω -peptides, mono-, di- and tri-substituted α glycopoly-peptides, polyamides, vinylogous sulfonamide peptide, ≘

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polyesters, polysaccharides, polycarbamates, polycarbonates, polyureas, polysulfonamide, conjugated peptides comprising e.g. prosthetic groups, polypeptidy/phosphonates, polyurethanes, azatides, oligo N-substituted sulfides, polynucleotides, PNAs, LNAs, morpholinos, oligo pyrrolinone, polyacetates, polystyrenes, polyvinyl, lipids, phospholipids, glycolipids, polyethylene glycols (PEG), polyethylenes, polydisulfides, polyarylene polyoximes, polyimines, polyethylenelmines, polyimides, polyacetals, polycyclic compounds comprising e.g. aliphatic or aromatic cycles, glycines, polyethers, ethoxyformacetal oligomers, poly-thioethers,

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including polyheterocyclic compounds, proteoglycans, and polysiloxanes,

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- a linker separating the functional entity from the complementing element. i i
- element is selected from a nucleotide sequence, such as a sequence of from 1-4 The building block according to claim 314, wherein the complementing nucleotides, such as from 1-3 nucleotides, such as 2 nucleotides or 3 nucleotides. 315.

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The building block according to claim 314, wherein the functional entity substituted amino acids, vinylogous amino acids, N-substituted glycin derivatives is selected from a precursor of an amino acid selected from alfa amino acids, beta amino acids, gamma amino acids, di-substituted amino acids, polyand other modified amino acids. 316.

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Figure 1. Chemical Display - Principle.

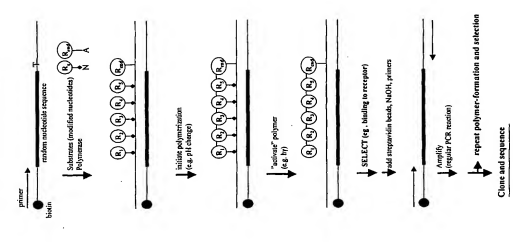


Figure 2a. An expanded set of base pairs.

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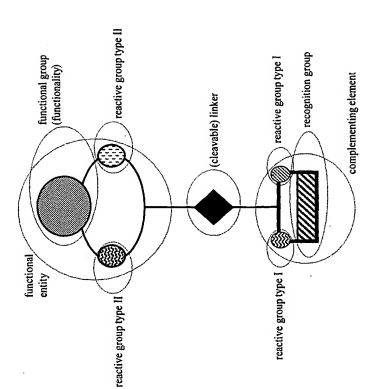
Figure 2b. An expanded set of base pairs.

Acceptor

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Donor Donor

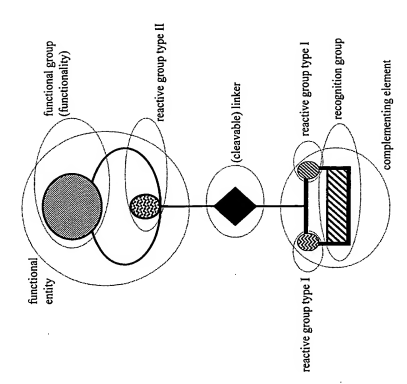
Figure 3. A monomer building block.



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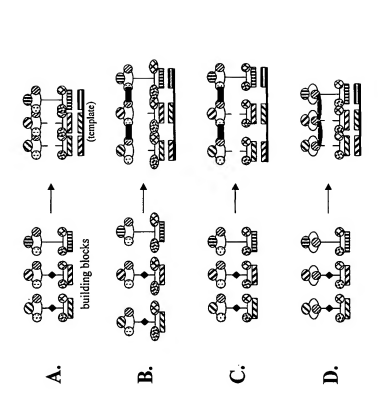
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Figure 4. A monomer building block with only one reactive group type II.



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Figure 5. Building blocks and the polymers resulting from template directed incorporation of the building blocks and their polymerization and activation.



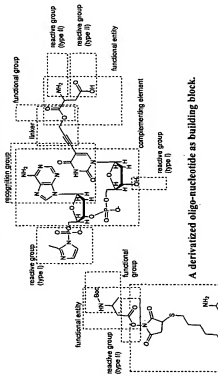
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Figure 6.

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A derivatized nucleotide as building block. functional entity reactive group (type II) reactive group (type I)

A derivatized di-nucleotide as building block. reactive group (type I)



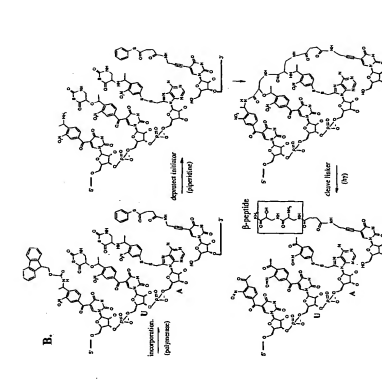
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Figure 7. C-terminal tagging of a β -peptide - incorporation, polymerization and activation, an example

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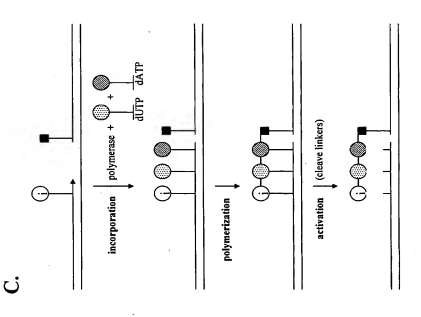
Figure 7. C-terminal tagging of a β -dipeptideincorporation, polymerization and activation. incorporation polymerase + + + activation | (cleave linkers) polymerization ن



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Figure 8. N-teminal tagging of a $\beta\text{-dipeptide}$ incorporation, polymerization and activation.



Bases and site of modification (-R)

Incorporated and extended nucleotide-derivatives (-R)

Nucleotide-derivatives for encoded polymer formation (-R)

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Figure 10. Cleavable linkers and protection groups,

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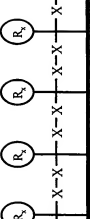
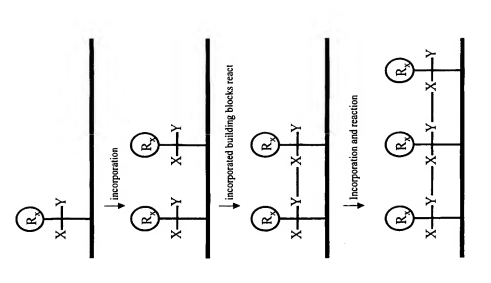


Figure 11, ex.1. Coumarin-based polymerization.

Figure 12. Polymerization between neighboring non-identical reactive groups type II.



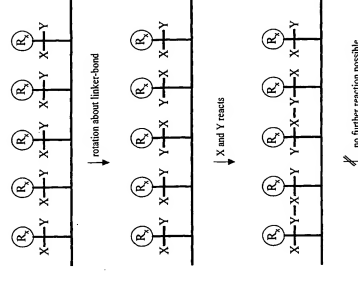
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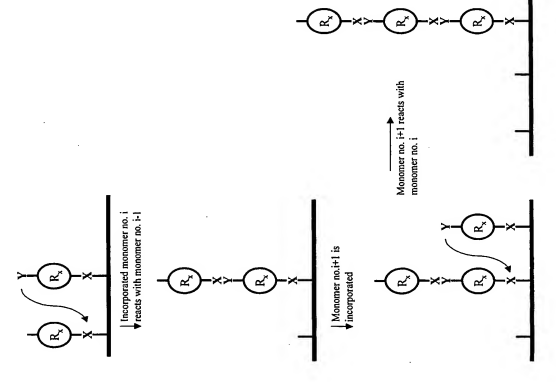
Figure 13. Cluster formation in the absence of directional polymerization.



no further reaction possible

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Figure 14. Zipping-polymerization and simultaneous activation.



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Figure 14, example 1. Polymerization and activation (thioesters)

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Increasing reactivity of thioester

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Figure 15. "Fill-in" polymerization (symmetric XX monomers).







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Figure 15, example 1. Poty-imine formation by fill-in polymerization.

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activate/deprotect

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Figure 15, example 3. Polyurea formation

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Figure 15, example 4. Chiral and achiral polyamide backbone formation

Figure 15, example 5. Polyphosphodiester formation.

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Figure 15, example 6. Polyphosphodiester formation with one reactive group in each monomer building

activate

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Figure 15, example 7. Pericyclic, "fill-in" polymerization.

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Figure 16. Encoded fill-in

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Figure 17. "Fill-in" polymerization (asymmetric XS monomers).

Figure 17, example 1. Fill-in polmerization by ketone-hydrazide reaction and by modified Staudinger ligation

Figure 18. "Zipping" polymerization.

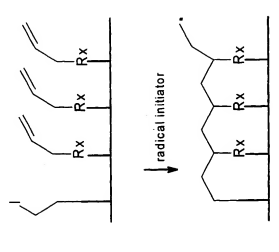
initiator

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Figure 18, example 1. Radical polymerization



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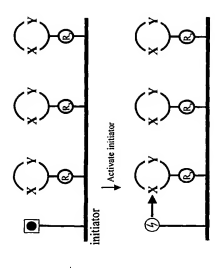
activate

Reaction activates reactive group Y

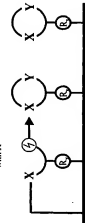
Initiator attacks neighboring X

Figure 18, example 2. Cationic polymerization

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Initiator reacts with neighboring reactive group x, resulting in ring-opening and activation of Y for reaction with X



Activated Y reacts with X in neighboring ring-structure

activation

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Figure 19, example 1. "Zipping" polymerization of N-thiocarboxyanhydrides, to form \(\beta\)-peptides.

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Figure 19, example 2. "Zipping" polymerization of 2,2-diphenylthiazinanone units to form β -peptides.

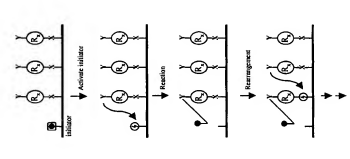
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Figure 20. Zipping-polymerization and activation by rearrangement.

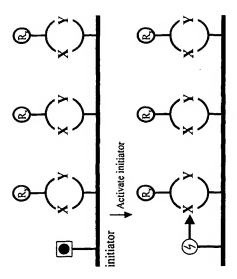


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Figure 21. Zipping-polymerization and activation by ring opening.



Initiator and X reacts, resulting in ring-opening and activation of Y. The functional entity is wimulaneously released from complementing element

Polymerisation and linker cleavage migrates along the template

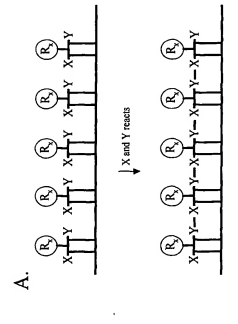
$$\mathbb{S} \left(\begin{array}{c} x \\ x \\ \end{array} \right)$$

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Figure 22. Directional polymer formation using fixed functional units.



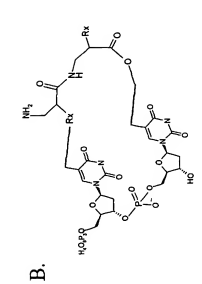


Figure 24.

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Figure 23.

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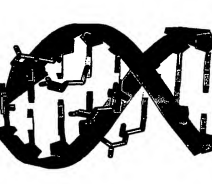
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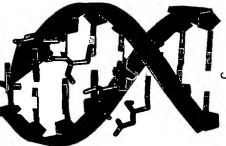
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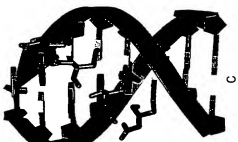
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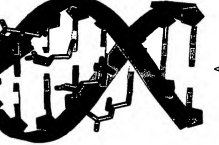


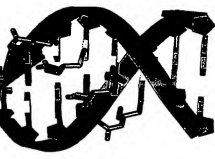


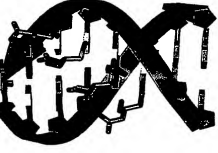




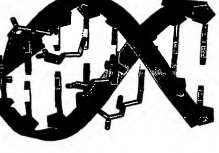




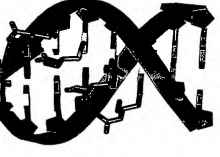




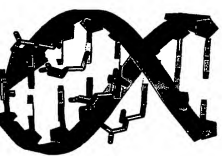




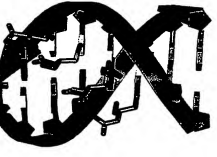






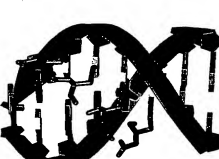


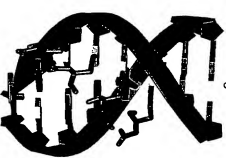




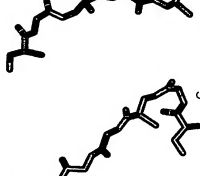


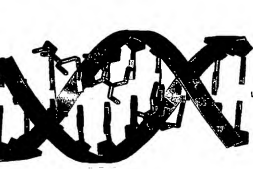


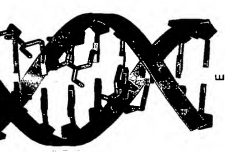












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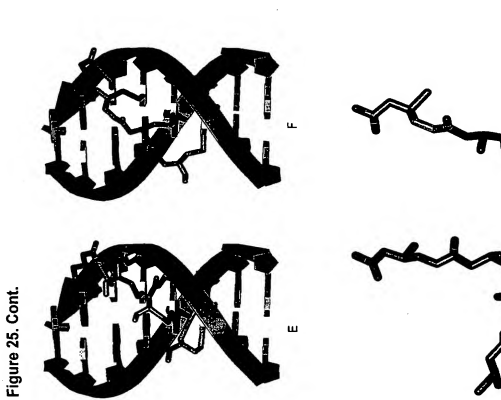
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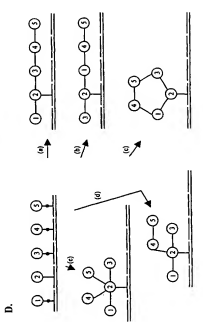
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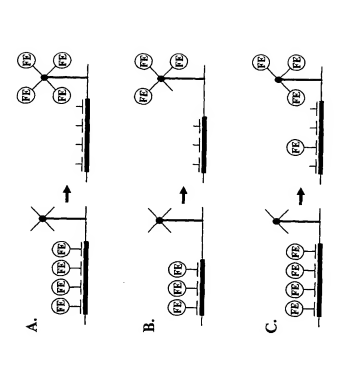
(FE) + (FE) Incorporation **↓**Cleavage Reaction Figure 26. Chemical Display - principle. læ (FE) Monomer building block with cleavable linker Monomer building block without cleavable linker ١Ą



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to, or lower than the number of reactive groups (Y). When the number of (X) and (Y) are different, scrambling results. Figure 27. The number of reactive groups (X) can be higher than, equal

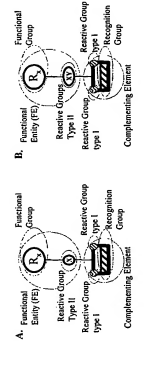


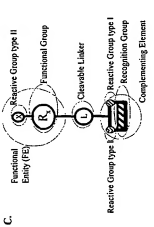
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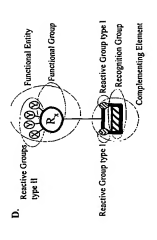
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Figure 28. Monomer building blocks.







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Figure 29. Templating involving simultaneous reaction and activation.

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Fig 30. Reaction types allowing simultaneous reaction and activation.

Nucleophilic substitution using activation of electrophiles

A. Acylating monomer building blocks - principle

C. Acylation Pyrazolone formation by reaction of hydrazines with $\beta\text{--}Ketoesters$

D. Acylation Isomation by reaction of hydroxylamines with $\beta\text{--Ketoesters}$

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E. Acylation

Pyrimidine formation by reaction of thioureas with \beta-Ketoesters

F. Acylation Pyrimidine formation by reaction of ureas with Malonates

G. Acylation Coumarine or quinolinon formation by a Heck reaction followed by a nucleophilic substitution



X' = Halogen, OTf, OMs Z = O, NH S'0 = X

H. Acylation Phthalhydrazide formation by reaction of Hydrazines and Phthalimides

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I. Acylation Diketopiperazine formation by reaction of Amino Acid Esters

J. Acylation Hydantoin formation by reaction of Urea and $\alpha\textsc{-substituted}$ Esters

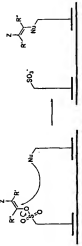
X = O, S X' = Hal, OTos, OMs, etc.

K. Alkylating monomer building blocks - principle Alkylated compounds by reaction of Sulfonates with Nucleofiles



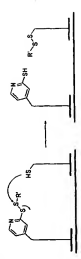
Nu = Oxygen-, Nitrogen-, Sufur- and Carbon Nucleophiles

L. Vinylating monomer building blocks - principle



Z = CN, COOR, COR, NO₂, SO₂R, S(=O)R, SO₂NR₂, F Nu = Oxygen- , Nitrogen- , Sultur- and Carbon Nucleophiles

M. Heteroatom electrophiles Disulfide formation by reaction of Pyridyl disulfide with Mercaptanes



N. Acylation Benzodiazepinone formation by reaction of Amino Acid Esters and Amino Ketones

Addition to carbon-hetero multiple bonds

O. Wittig/Horner-Wittig-Emmons reagents Substituted alkene formation by reaction of Phosphonates with Aldebydes or Ketones

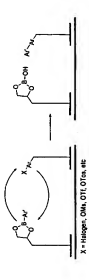
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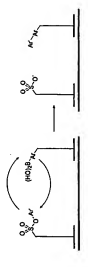
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Transition metal catalysed reactions

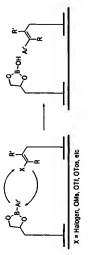
P. Arylation Biaryl formation by the reaction of Boronates with Aryls or Heteroaryls



Q. Arylation Biaryl formation by the reaction of Boronates with Aryls or Heteroaryls



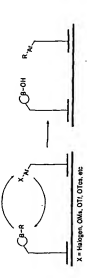
R. Arylation Vinylarene formation by the reaction of alkenes with Aryls or Heteroaryls



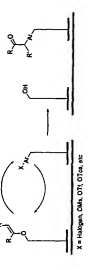
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S. Alkylation Alkylation of arenes/hetarens by the reaction with Alkyl boronates

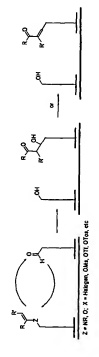


T. Alkylation Alkylation of arenas/hetarenes by reaction with enolethers



Nucleophilic substitution using activation of nucleophiles

U. Condensations Alkylation of aldchydes with enolethers or enamines



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V. Alkylation Alkylation of aliphatic halides or tosylates with enolethers or enamines

Cycloadditions

W. [2+4] Cycloadditions

X. [2+4] Cycloadditions

Y, CN, COOR, COR, NO2, SO2R, S(=O)R, SO2NR2, F

Y. [3+2] Cycloadditions

Y, CN, COOR, COR, NO2, SO2R, S(=0)R, SO2NR2, F

Y, CN, COOR, COR, NO2, SO2R, S(=0)R, SO2NR2, F Z. [3+2] Cycloadditions

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Figure 31. Templating involving non-simultaneous reaction and activation.

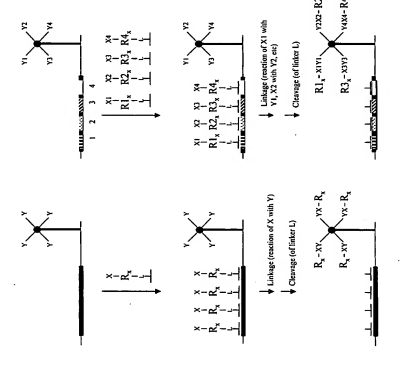


Figure 32. Pairs of reactive groups X,Y and the resulting bond XY. Nucleophilic substitution reaction

Aromatic nucleophilic substitution

SUBSTITUTED AROMATIC COMPOUNDS

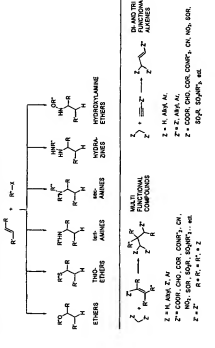
Transition metal catalysed reactions

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Addition to carbon-carbon multiplebonds



Cycloaddition to multiple bounds

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Addition to carbon-hetero multiple bonds

2 = COOR, CHO, COR, SOR, SO2R, CN, NO2, ect.

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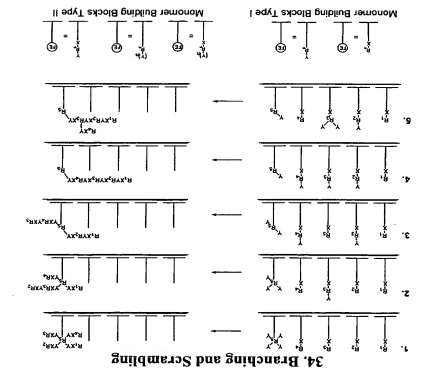
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33. Anchorage - Examples

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35. Monomer Building Blocks - Examples of Linker Design







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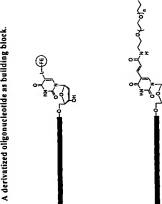
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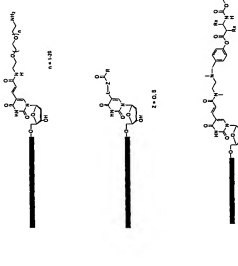
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36. Formation of Monomer Building Blocks – Examples

Figure 38. An oligonucleotide-based monomer building block. Example of complementing element design, allowing for high monomer diversity.

0X 4 B0X 5 B0X 6	monomer diversity 1024 1024 1024 1024 4 4	
BOX 1 BOX 2 BOX 3 BOX 4 BOX 5 BOX 6	sequence XXXXXATTTTXXXXXX XXXATTTTAXXXXXXX XXXATTTTAXXXXXXXX	
BOX	BOX 1 2 2 3 3 4 4 4 4 5 5 6 6	
⋖		

X = G or C

Coding Element (BOX 1) GCGCGATATTTGGGCC Complementing Element CGCGCTATAAACCCGG ä

Coding Element (BOX 6) GAGAGTTCTTCGCGGG Complementing Element CTCTCAAGAAGCGCCC

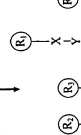
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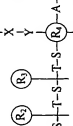
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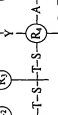
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Figure 38, continued.

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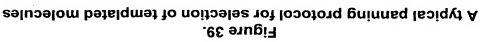


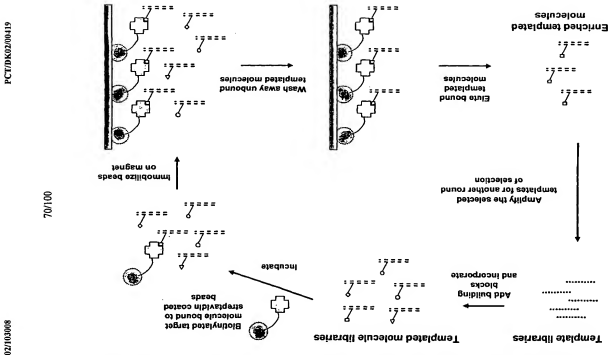




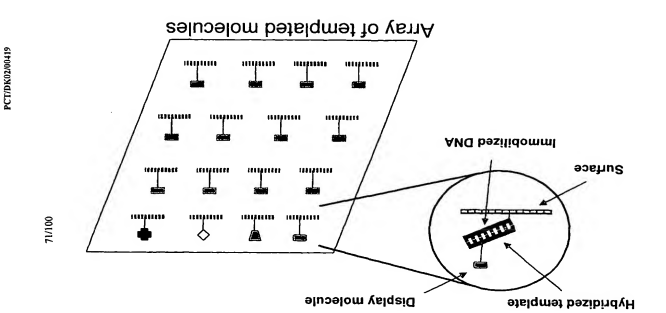
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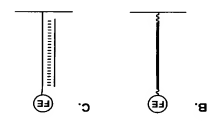




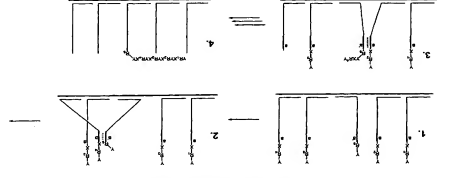
Array of templated molecules Figure 40



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42. Use of Zipper Box



B. Example of a building block in which the zipper box consists of an oligonucleotide sequence:

Z= Amino-Modifier Glen C6 dT 10-1039-X= Carboxy-dT Glen cat.no. 10-1035-

S'-CGACCTCTGGATTGCATGGTCATGGCTGACTGTCCGTCGAATGTGTCCAGTTACX
Annealing region Linker region Zipper region Annealing region logilO

Oligo2

5'-ZGTAACACCTGTGTAAGCTGCCTGTCAGTCGGTACTGACCTGTCGAGCATCCAGCT

Zipper region Annealing region

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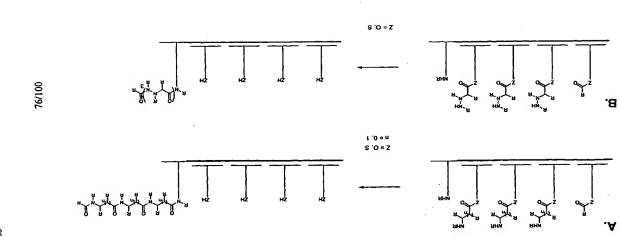
Figure 43. Templated synthesis of organic compounds - examples.

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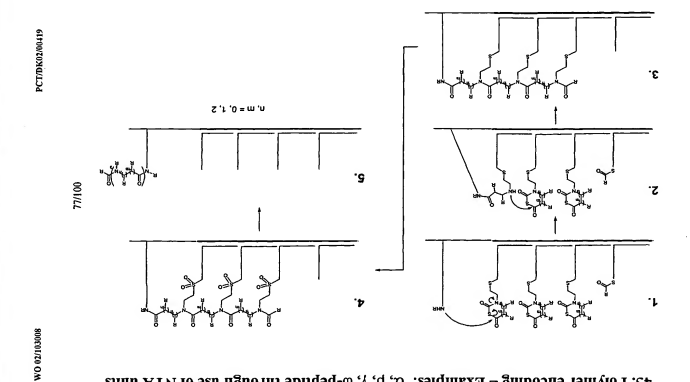
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44. Polymer encoding – Examples: peptide (α , β), peptoid or hydrazino peptide



45. Polymer encoding – Examples: α , β , γ , ω -peptide through use of NTA units



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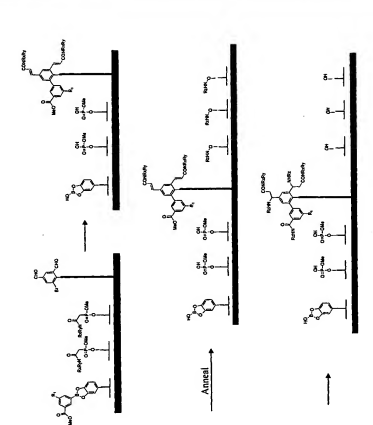
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46. Example 1: Double templating



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A. Linker for the formation of Ketones, Aldehydes, Amides and Acids

Figure 47. Cleavable Linkers

. Linker for the formation of Ketones, Amides and Acids

). Linker for the formation of Aldehydes and Ketones

). Linker for the formation of Alcohols and Acids

E. Linker for the formation of Amines and Alcohols
$$\mathbb{R} \underbrace{X_0 \int_{W^{-R}}^{R} \frac{1}{W^{-R}}}_{R} \underbrace{X_{OH} \cdot R^{-Net}}_{R}$$

F. Linker for the formation of Esters, Thioesters , Amides and Alcohols

G. Linker for the formation of Sulfonamides and Alcohols

H. Linker for the formation of Ketones, Amines and Alcohols

1. Linker for the formation of Ketones, Amines, Alcohols and Mercaptunes

. Linker for the formation of Biaryl and Bihetaryl

K. Linker for the formation of Benzyles, Amines, Anilins Alcohols and Phenoles

M. Linker for the formation of Glycosides S-R TCEP R-SH + R-SH H_LO / Dictains TCEP = Intq2-carboxysthyphotophie

i. Linker for the formation of Aldehydes and Glyoxylamide:

D. Linker for the formation of Aldchydes, Ketones and Aminoalcohols

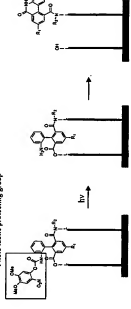
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Figure 48. Post-templating modification oftemplated molecule

A Rearrangement and cleavage in one step, eg:

Photo labile protecting group

42



B Reaction of functional groups present in a templated molecule

B1 Intramolecular Michael addition:

B2 Intermolecular Michael addition:

B3 Reaction of phenylenediamines and aldehydes to form benzimidazoles:

B4 Reduction of multiple bonds:

9 2 8 6 10 11 15 13 14 18

S

Figure 49

Start-

C Post templating modification of linker to extend the spacing between the template and the templated molecule.

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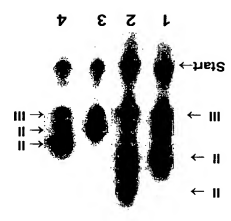
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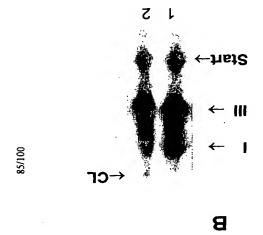
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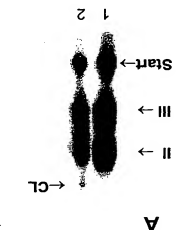
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Figure 53

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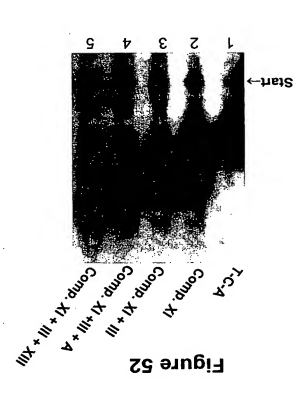
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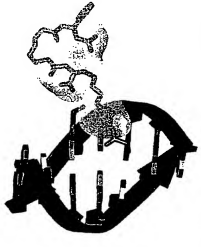
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Сотр.

Figure 55A. DNA template with three modified nucleotides (Compound V activated by NHS) incorporated (all shown in stick models coloured by residue)

between compound X and compound V and 2) bridging of compound X and compound II using added reagent BS₃ Figure 55B. DNA template bearing modifications after 1) reaction



ε

7

Figure 54

9

6

← CF

8 L 9 4

Figure 55C. DNA displaying the templated molecule after hydrolysis of the two hydrolyzable ester elements. All modifications are shown in stick representation and in addition the end product is highlighted by its surface.

All figures are created using Accelrys' ViewerLite 4.2 with structures calculated using Schrödinger's MacroModel 7.2

8£4A -\+

AY2 .gi7

4\-Ab38

Incubation temperatur

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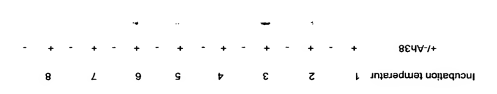
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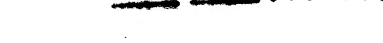




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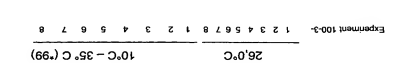


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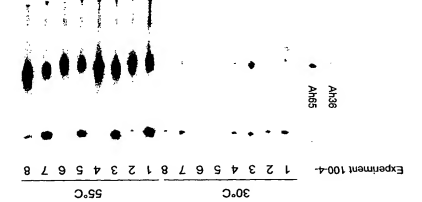




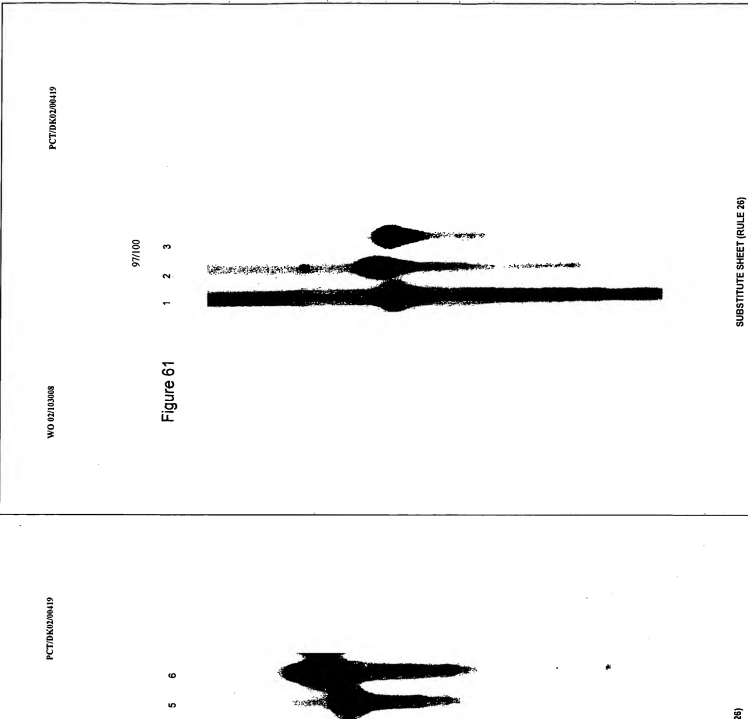
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Figure 60

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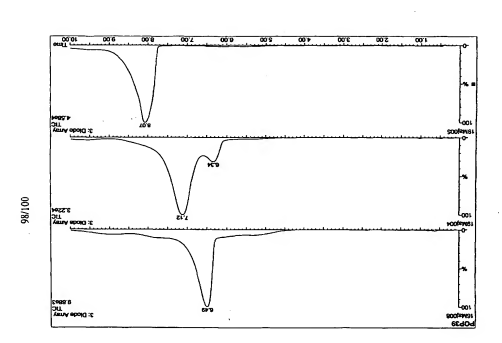
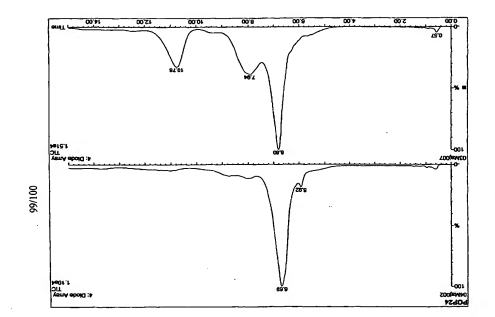


Figure 63



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AH17+AH18+T4 ligase AH19+AH20+ T4 ligase

AH19 AH 17

CI2N 15/10, (21) International Application Number: PCI/DK02/00419 (22) International Filing Date: 20 June 2002 (20,06,2002) (51) International Patent Classification*: C12P 1/00, 19/34, C0711 21/00, C12P 21/02 (43) International Publication Date 27 December 2002 (27.12.2002) International Bureau

PCT (19) World Intellectual Property Organization

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DX DX

20 June 2001 (20.06.2001) 15 March 2002 (15.03.2002)

(30) Priority Data: PA 2001 00962 PA 2002 00415

(71) Applicant (for all designated States except US): NUEVO-LUTION A/S [DK/DK]; Rønnegade 8, 5th floor, DK-2100

Copenhagen (JUK).

[Continued on next page]

(54) Title: TEMPLATED MOLECULES AND MITHODS FOR USING SUCH MOLECULES

Chemical Display - Principle. <u>ත්ත්ත්ත්ත්</u>

@@@@@

(57) Abstract: The present invention relates to a method for synthesising templated molecules. In one aspect of the invention, the templated molecules are linked to the template which templated the synthesis thereof. The intion allows the generation of libraries which can be screened for e.g., therapeutic activity.

. repeat polymer-formation and selection 900000 and strepteridio beach. NaOH, prime SELECT (cg. bledling to recepto em(pd_spage_ba) WO 02/103008 A3

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Fig. 64

Lene [DKDK]; Solsikkemarken 21, DK-2830 Virum (8 (DK), NØRREGAARD-MADSEN, Mads [DKDK]; Bilebakken 5, DK-3460 Birkerad (DK), ANDERS GIOSKESEN, Michael [DKDK]; Plantagetrogen 8, DK-2950 Vedrek (DK), SCHRØDER GIAD, Sanne [DKDK]; Viggo Barfoods Alle 59, DK-2750 Ballerup

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Published:

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	INTERNATIONAL SEARCH REPORT	Inte nat Application No PCT/DK 02/00419	
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INTERNATIONAL SEARCH REPORT

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vational application No. PCT/DK 02/00419

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)	ole (Continuation of item 1 of first sheet)	
This international Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	daims under Article 17(2)(a) for the following reasons:	
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:	is Authority, namely:	
 Claims Nos.: because they refare to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: 	comply with the prescribed requirements to such bedifically:	
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a)	with the second and third sentences of Rule 6.4(a).	
3ox II Observations where unity of invention is tacking (Continuation of Itam 2 of first sheet)	tion of item 2 of first sheet)	
	ial application, as follows:	
see additional Sheet		
. Searchable claims.	this international Search Report covers all	
. As all soarchable claims could be searched without eflort justifying an additional fee, this Authority did not invite payment of any additional fee.	ddilonal fee, this Authority did not invite payment	
. The sony some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:	the applicant, this International Search Report.	
No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-232,236 (complete); 237-242,244-253,265-289,292,296-306 (partially)	naequenty, this International Search Report is laims Nos.: 1889,292,296-306 (partially)	
remark on Protest The additional search less were accomp	The additional search leas were accompanied by the applicant's protest. No protest accompanied the payment of additional search leas.	

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FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 219

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

 Claims: 1-232, 236, (237-242, 244-253, 265-289, 292, 296-306)-partially A method for synthesising a template molecule comprising a steps of illurality of functional groups, said method comprising the steps of i) providing at least one template comprising a sequence of n coding elements, wherein each coding element comprises at least one recognition group capable of recognising a predetermined complementing element, and wherein n is an integer of more than 1, ii) providing a plurality of building blocks, wherein each building block comprises a) at least one complementing element comprising at least one recognition group capable of recognising a predetermined coding element, b) at least one functional group and at least one reactive group, and c) at least one linker separating the at least one functional entity from the at least one ecomplementing element, ii) contacting each of said coding elements with a complementing element, ii) contacting each of said coding elements, ii) contacting each of said coding elements of a reaction involving reactive groups, a linking, by means of a reaction involving reactive groups, a functional group of antlest one functional entity to a functional group of antlest the complementing template of being linked by means of a linker to the complementing template or template that template the synthesis of the templated molecule does not involve ribosome mediated translation of a nucleic acid:

 Claims: 233, (237-242, 244-253, 265-289, 292, 296-306)-partially A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule does not comprise or consist of an a-peptide or a nucleotide;

3. Claimš: 234, (237-242, 244-253, 265-289, 292, 292)

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule does not comprise or consist of a monosubstituted a-peptide or a nucleotide;

4. Claims: 235, (237-242, 244-253, 265-289, 292. 296-306)-partially A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule does not comprise or consist of a peptide or a nucleotide;

5. Claim : 243

A complex comprising a template molecule and the template that template the synthesis of the template molecule;

6. Claims: 254, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template molecule does not comprise or consist of an alpha-peptide;

7. Claims: 255, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the templated molecule, wherein the template molecule and experient the template molecule and the synthesis of the templated appendix that template molecule are not comprise a monosubstituted a-peptide;

8. Claims: 256, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the templated molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template molecule, one an arrest of a arrest of a

9. Claims: 257, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template is not a natural nucleotide, when the template molecule is an a-peptide;

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10. Claims: 258, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template does not consist exclusively of natural nucleotides, when the template molecule is a peptide comprising exclusively monosubstituted a-amino

11. Claims: 259, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template is not a natural nucleotide, when the template molecule is a natural apportide;

12. Claims: 260, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template is not a nucleotide, when the template a natural a-peptide;

13. Claims: 261, (265-289, 292, 296-306)-partially

A templated molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template is not a nucleotide, when the template molecule is a monosubstituted a-peptide;

14. Claims: 262, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template is not a nucleotide, when the template is an a-peptide;

15. Claims: 263, (265-289, 292, 296-306)-partially

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A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template is not a natural nucleotide, when the template molecule; as not a natural nucleotide, when the template molecule is a peptide;

16. Claims: 264, (265-289, 292, 296-306)-partially

A template molecule comprising a sequence of covalently linked, functional groups, wherein the template molecule is linked by means of a linker to the complementing template or template that template the synthesis of the template molecule, wherein the template is not a nucleotide, when the template molecule is a peptide;

17. Claims: 290, (292, 296-306)-partially

A molecule comprising a sequence of covalently linked building blocks, wherein the sequence of covalently linked building blocks comprises a sequence of complementing elements forming a complementing template capable of complementing the template that template the synthesis of the template molecule, and wherein the template molecule is linked to the complementing template or template that template its synthesis;

18. Claims: 291, (292)-partially

A templated molecule according to any of the previous claims, wherein the templated molecule comprises a sequence of functional entities comprising at least one functional group, and optionally at least one reactive group type 11, and wherein each functional entity is linked to a compinementing element or a template that template the synthesis of the templated molecule;

19. Claim : 293

A method for screening template molecules potentially having a predetermined activity, said method comprising the step of providing a target molecule or a target entity, including a surface, and obtaining template molecules having an affinity for-or an effect on-said target molecule or target entity;

20. Claim : 294

A method for assaying an activity potentially associated with a template molecules, said method comprising the step of providing a target molecule or a target entity, including

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a surface, and obtaining template molecules having an affinity for-or an effect on-said target molecule or target entity, and determining the activity of the templated molecule;

21. Claim : 295

A method for selecting complexes or template molecules having a predetermined activity, said method comprising the step of performing a selection procedure and selecting templated molecules based on predetermined selection criteria:

22. Claim : 307

A method for amplifying the complementing template or the template that template the synthesis of the templated molecule having, or potentially having a predetermined activity, said method comprising the step of contacting the template with amplification means, and amplifying the template.

23. Claim: 308

A method for amplifying the complementing template or the template that template the synthesis of the templated molecule having, or potentially having, a predetermined activity, said method comprising the steps of i) contacting the template with amplification means, and amplifying the template, and ii) obtaining the templated molecule in an at least two-fold increased amount.

24. Claim: 309

A method for altering the sequence of a templated molecule, including generating a template molecule comprising a novel or altered sequence of functional groups, wherein said method preferably comprises the steps of i) providing a first complementing template or a first template capable of templating the first templates or first templates capable of templating a plurality of first templates capable of templating a plurality of first template molecules, ii) mutating or modifying the sequence of the first complementing template or the first template, or the plurality of first complementing templates or first templates or second complementing templates, wherein said second templates or second complementing templates, wherein said second template (s) or complementing templates, wherein said second template (s) or complementing templates, wherein said second template is or a template ing the synthesis of a second template molecule, or a

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plurality of second template molecules, wherein said second template molecule (s) comprises a sequence of covalently linked, functional groups that is not identical to the sequence of functional groups of the first template molecule(s), and optionally iii) templating by means of said second template(s) or complementing tem plate (s) a second template molecule, or a plurality of such second tem plated

25. Claims: 310-313

A method for altering the sequence of a template molecule, including generating a template molecule comprising a novel or altered sequence of functional groups, wherein said method preferably comprises the steps of i) providing a plurality of first complementing templates or first template molecules, ii) recombining the sequences of the plurality of first complementing templates or first templates, and generating a second template or a second complementing template, or a plurality of second templates or second complementing templates, wherein said second template(s) or complementing templates, wherein said second template(s) or complementing templates, wherein said second template molecule(s) comprises a sequence of covalently linked, functional groups that is not identical to the sequence of functional groups of the first template molecule(s), and optionally iii) templating by means of second template molecules, or a plurality of such second template molecules, second template molecules, second template molecules, or a plurality of such second template molecules.

26. Claims: 314-316

A building block comprising i) a complementing element capable of specifically recognising a coding element having a recognition group, said complementing element being selected from nucleotides, amino acids, antibodies, antigens, proteins, peptides, and molecules with nucleotide recognizing ability, ii) at least one functional entity selected from a precursor of a-peptides, p-peptides, y-peptides, w-peptides, p-peptides, polyamides, vinylogous polypeptides, glycopoly-peptides, polyamides, vinylogous sulfonamide peptide, polysulfonamide, conjugated peptides comprising e. g. prosthetic groups, polyareters, polyachenies, polyarethanes, azatides, oligo N-substituted glycines, polyarethanes, azatides, oligo M-substituted glycines, polyathylene glycols (PEG), polyethylenes, polyatides, polyarylene

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sulfides, polynucleotides, PNAs, LNAs, morpholinos, oligo pyrrolinone, polyoximes, polyimines, polyethyleneimines, polyimides, polyacetals, polyacetates, polystyrenes, polyimides, pipospholipids, glycolipids, polycylic compounds comprising e. g. aliphatic or aromatic cycles, including polyheterocyclic compounds, proteoglycans, and polysiloxanes, and iii) a linker separating the functional entity from the complementing element:

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